

**SQUIRE'S
COMPANION**

TO THE LATEST EDITION OF THE

BRITISH PHARMACOPŒIA.

EIGHTEENTH EDITION

**SQUIRE'S
COMPANION
TO THE LATEST EDITION OF THE
BRITISH PHARMACOPŒIA**

**COMPARING THE STRENGTH OF ITS VARIOUS PREPARATIONS
WITH THOSE OF THE
UNITED STATES, AND OTHER FOREIGN PHARMACOPŒIAS,
TO WHICH ARE ADDED
NOT OFFICIAL PREPARATIONS,
AND
PRACTICAL HINTS ON PRESCRIBING**

**BY
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Eighteenth Edition.

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The Board of Inland Revenue have consented to recognise 'Squire's Companion to the British Pharmacopœia' as a 'well known' book of reference in connection with the exemption from Medicine Stamp Duty in favour of 'known, admitted and approved' remedies

PREFACE

TO EIGHTEENTH EDITION.

THE SEVENTEENTH EDITION of SQUIRE'S COMPANION was published in 1899, and since that time a great advance has been made in the testing of Drugs and Chemicals used in medicine, a very large number of new synthetic products have also been introduced. Several foreign Pharmacopœias have issued new editions, viz, Austrian, Belgian, Danish, Dutch, French, German, Italian, Russian, Spanish, Swedish, Swiss and United States, the Japanese is a new Pharmacopœia.

To bring SQUIRE'S COMPANION up to date and to make it conform to modern requirements, it has been necessary to practically re-write it from cover to cover. The general arrangement of the book remains the same as before. Substances which are official in the British Pharmacopœia have the names in larger type than those which are 'Not Official,' and the same distinction also applies to the preparations, it is therefore quite easy to see at a glance whether any particular substance or preparation is official or not. This is, moreover, supplemented by a list of 'Official' and 'Not Official' preparations given under each substance immediately following the dose, so that a prescriber can quickly ascertain the various forms in which any medicament can be given. Following the precedent of the previous editions, the formulas are given in parts, solids by weight, liquids by measure, and where it has been necessary to depart from this course, it is stated in the text. In dealing with German and other Continental Pharmacopœias, it must be understood that parts refer to parts by weight.

The arrangement of the matter and the headings to the several paragraphs, which have always been the distinctive feature of the COMPANION, have been retained. 'Solubility,' 'Medicinal Properties,' 'Dose,' 'Prescribing Notes,' 'Incompatibles,' list of 'Official Preparations,' 'Antidotes,' and 'Foreign Pharmacopœias,' this arrangement having been found very convenient to those who use the work as a book of reference. The term 'Medicinal Properties,' although very old fashioned, has been retained in order to keep the plan of the book uniform with previous editions, this portion has been carefully read and corrected by Dr. Taylor Grant.

The 'Descriptive Notes' have been written specially for this book by Mr. E. M. Holmes, they deal with the principal distinguishing features of the vegetable drugs, the commercial qualities, and the probable contaminations and sophistications. He has also assisted in the revision of the Organic Materia Medica.

The 'Tests' have been entirely re-written, and include a critical comparison of the tests given in the Pharmacopœias of Great Britain,

Germany, and the United States of America, together with such further tests as have been found useful in the laboratory of the Author. The French Codex, so recently published, could not be included in the critical comparison mentioned above, but the more important tests therein contained have been dealt with, and this work has delayed the publication of the COMPARISON beyond the time at which it was intended to have been ready.

The standardisation of preparations is exciting special interest, and considerable attention has been devoted to this subject, the standards and methods for ensuring them, which are employed in various Pharmacopœias are given, and are supplemented by figures which have been obtained from work devoted to this purpose in the author's laboratory.

A large number of formulas which appear in the British Pharmaceutical Codex are compared with those previously published in books of reference in common use.

The chapter on Therapeutic Agents of Microbial Origin has been revised and partly re-written by Dr. R. Tanner Hewlett.

The author wishes to acknowledge his indebtedness to the members of his staff who have assisted in the preparation of this edition, and to various friends who have supplied information on subjects with which their names have been particularly associated. His thanks are specially due to Mr Charles M. Caines, who has devoted the whole of his attention to the Chemical portion, and who has been the principal chemist employed on the experimental work.

P. W. SQUIRE.

413 Oxford Street,
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ABBREVIATIONS.

<i>Allen</i>	-	Allen's Commercial Organic Analysis
<i>A J P.</i>	=	American Journal of Pharmacy
<i>B M J</i>	=	British Medical Journal
<i>B M J E</i>	=	British Medical Journal Epitome
<i>B P</i>	=	British Pharmacopoeia 1884
<i>B P.O. Formulary</i>	=	British Pharmaceutical Conference (Unofficial Formulary, 1901)
<i>B P C.</i>	=	British Pharmaceutical Codex (1907) and Corrections
<i>Brompton</i>	=	Pharmacopoeia of the Brompton Hospital for Consumption, 1891
<i>Brussels Conference</i>	=	International Agreement respecting the Unification of the Pharmacopoeial Formulas for Potent Drugs (Signed at Brussels, November 21, 1906)
<i>Charing Cross</i>	=	Pharmacopoeia of the Charing Cross Hospital, 1901
<i>C F</i>	=	Canadian Formulary of Unofficial Preparations
<i>C D</i>	=	Chemist and Druggist
<i>Central Throat</i>	=	Pharmacopoeia of the Central London Throat and Ear Hospital, 1901
<i>City of London Chest</i>	=	Pharmacopoeia of the City of London Hospital for Diseases of the Chest, 1901
<i>East London</i>	=	Pharmacopoeia of the East London Hospital for Children, 1903
<i>Exelina</i>	=	Pharmacopoeia of the Exelina Hospital for Sick Children, 1906
<i>F.T</i>	=	Folia Therapeutica
<i>Great Northern</i>	=	Pharmacopoeia of the Great Northern Central Hospital, 1899
<i>G P.</i>	=	General Practitioner
<i>Guy's</i>	=	Pharmacopoeia of Guy's Hospital, 1891
<i>Hager</i>	=	Hager's Handbuch der Pharmaceutischen Praxis.
<i>Ind. and Col. Add</i>	=	Indian and Colonial Addendum to the British Pharmacopoeia
<i>I.M.G.</i>	=	Indian Medical Gazette.
<i>I.M.R.</i>	=	Indian Medical Record
<i>J.C.S. Abs.</i>	=	Journal of the Chemical Society Abstracts.
<i>J.C.S. Trans</i>	=	Journal of the Chemical Society Transactions.
<i>J.C.S.I.</i>	=	Journal of the Society of Chemical Industry
<i>King's</i>	=	Pharmacopoeia of King's College Hospital, 1901.
<i>L.</i>	=	Lancet.
<i>Lock</i>	=	Pharmacopoeia of the London Lock Hospital, 1896.
<i>London</i>	=	Pharmacopoeia of the London Hospital, 1909.
<i>L.M.R.</i>	=	London Medical Recorder
<i>London Ophthalmic</i>	=	Pharmacopoeia of the Royal London Ophthalmic Hospital (late Moorfields), 1901
<i>London Skin</i>	=	Pharmacopoeia of the London Skin Hospital, 1903.
<i>Martindale</i>	=	Martindale and Westcott's Extra Pharmacopoeia.
<i>M.A.</i>	=	Medical Annual
<i>M.P.</i>	=	Medical Press and Circular
<i>M.T.</i>	=	Medical Times and Gazette.
<i>Marck</i>	=	Marck's Archives
<i>Middlesex</i>	=	Pharmacopoeia of the Middlesex Hospital, 1899.
<i>Murrell</i>	=	What to do in Cases of Poisoning (Murrell).
<i>Edin. Pharm.</i>	=	Edinburgh Pharmacopoeia.
<i>Pharm. Form.</i>	=	Pharmaceutical Formulas (Peter MacEwan)
	=	Pharmacopoeia Germanica (Editio IV.).
	=	Pharmaceutical Journal (Second, Third, and Fourth Series).
	=	Pharmaceutical Journal Formulary.

<i>Ph Lond</i>	=	Pharmacopœia Londinensis, 1851
<i>P R</i>	=	Pharmaceutical Record (New York)
<i>Pr</i>	=	Practitioner
<i>Proc Amer Pharm Assoc</i>	=	Proceedings of the American Pharmaceutical Association
<i>Ringer</i>	=	Ringer's Handbook of Therapeutics
<i>Royal Chest</i>	=	Pharmacopœia of the Royal Hospital for Diseases of the Chest, 1891
<i>R D H</i>	=	Pharmacopœia of the Royal Dental Hospital, 1907
<i>Royal Free</i>	=	Pharmacopœia of the Royal Free Hospital, 1904
<i>Samaritan</i>	=	Pharmacopœia of the Samaritan Free Hospital, 1906
<i>St Bartholomew's</i>	=	Pharmacopœia of St Bartholomew's Hospital, 1900
<i>St George's</i>	=	Pharmacopœia of St George's Hospital, 1907
<i>St John's</i>	=	Pharmacopœia of St John's Hospital for Diseases of the Skin, 1894
<i>St Mary's</i>	=	Pharmacopœia of St Mary's Hospital, 1904
<i>St Thomas's</i>	=	Pharmacopœia of St Thomas's Hospital, 1902
<i>Schimmel</i>	=	Schimmel's Semi-Annual Reports
<i>Sheffield Union</i>	=	Pharmacopœia of the Sheffield Union, 1903
<i>Sick Children (G O S)</i>	=	Pharmacopœia of the Hospital for Sick Children, Great Ormond St., 1900
<i>Squibb</i>	=	Squibb's Ephemeris
<i>T'G</i>	=	Therapeutic Gazette (Philadelphia)
<i>Throat</i>	=	Pharmacopœia of the Hospital for Diseases of the Throat, Golden Square, 1901
<i>U S P</i>	=	United States Pharmacopœia
<i>U S N F</i>	=	National Formulary of the American Pharmaceutical Association
<i>University</i>	=	Pharmacopœia of the University College Hospital, 1904
<i>University (1907)</i>	=	Pharmacopœia of the University College Hospital, 1907
<i>Victoria</i>	=	Pharmacopœia of the Victoria Hospital for Children, 1904
<i>Westminster</i>	=	Pharmacopœia of the Westminster Hospital, 1902
<i>Westminster Ophthalmic</i>	=	Pharmacopœia of the Royal Westminster Ophthalmic Hospital, 1901
<i>Women</i>	=	Pharmacopœia of the Hospital for Women, 1883
<i>Y B P</i>	=	Year book of Pharmacy

EXAMPLE *L '04*, i 56, refers to *Lancet*, 1904, Volume I, page 56

The British Pharmacopœia, published in 1898, is in this work compared with the latest editions of the foreign Pharmacopœias, which are as follows —

Austrian	published in 1906	Mexican	published in 1896
Belgian	" " 1906	Norwegian	" " 1895
Danish	" " 1907	Portuguese	" " 1876
Dutch	" " 1905	Russian	" " 1902
French	" " 1908	Spanish	" " 1905
German	" " 1900	Swedish	" " 1901
Hungarian	" " 1888	Swiss	" " 1907
Italian	" " 1902	United States	" " 1905
Japanese	" " 1907		

and are thus expressed — Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss, U.S.

The following works have also been consulted — 'United States Dispensatory', Dorvault, 'L'Officine', Gray's 'Supplement', Gildemeister and Hoffmann, 'Volatile Oils', Parry, 'Chemistry of Essential Oils', Sutton, 'Volumetric Analysis', Pluckiger and Hanbury, 'Pharmacographia', Vogl, 'Anatomischer Atlas zur Pharmakognosie', Vogl, 'Pharmakognosie', Planchon and Collin, 'Les Drogues Simples d'Origine Végétale', Tschirch and Oesterle, 'Anatomischer Atlas', Moeller, 'Leitfaden zu Microscopisch pharmacognostischen Übungen', Schneider, 'Powdered Vegetable Drugs', Greenish and Collin, 'An Anatomical Atlas of Vegetable Powders', Koch, 'Die Microscopische Analyse der Drogenpulver.'

SYMBOLS, AND ATOMIC WEIGHTS OF THE ELEMENTARY BODIES, MENTIONED IN THE BRITISH PHARMACOPOEIA

H = 1.00

Elementary Bodies	Symbol and Atomic Weights	International Atomic Weights H = 1
Aluminium	Al = 26.90	26.90
Antimony (Stibium)	Sb = 119.00	119.00
Arsenium	As = 74.50	74.40
Barium	Ba = 136.40	136.40
Bismuth	Bi = 207.30	206.90
Boron	B = 10.85	10.90
Bromine	Br = 79.35	79.36
Calcium	Ca = 89.71	89.80
Carbon	C = 11.91	11.91
Cerium	Ce = 139.20	139.20
Chlorine	Cl = 35.19	35.18
Chromium	Cr = 51.71	51.70
Copper (Cuprum)	Cu = 63.12	63.10
Gold (Aurum)	Au = 195.70	195.70
Hydrogen	H = 1.00	1.00
Iodine	I = 125.90	125.90
Iron (Ferrum)	Fe = 55.60	55.60
Lead (Plumbum)	Pb = 205.35	205.35
Lithium	Li = 6.97	6.98
Magnesium	Mg = 24.18	24.18
Manganese	Mn = 54.52	54.60
Mercury (Hydrargyrum)	Hg = 198.80	198.50
Nitrogen	N = 13.91	13.98
Oxygen	O = 15.84	15.88
Phosphorus	P = 30.80	30.77
Platinum	Pt = 193.30	193.80
Potassium (Kalium)	K = 38.83	38.86
Silver (Argentum)	Ag = 107.11	107.12
Sodium (Natrium)	Na = 22.88	22.89
Sulphur	S = 31.82	31.88
Stannum	Sn = 118.20	118.10
Zinc	Zn = 64.91	64.90

THE WEIGHTS AND MEASURES OF THE BRITISH PHARMACOPŒIA
AT THE TEMPERATURE OF 60° FAHRENHIT

WEIGHTS

The Avoirdupois pound = 16 oz = 7000 grains
 1 oz = 437 5 grains
 1 gr = 1 grain

In addition to the use of the Imperial weights it is permitted in the Act of 1878 that drugs when sold by retail may be sold by apothecaries' weight. The use in trade of a weight or measure of the metric system was made lawful by the Weights and Measures (Metric System) Act, 1897.

The Preface to the British Pharmacopœia states 'It is strongly urged upon all medical men to avoid the use of the terms ounce and pound with reference to any other than the avoirdupois or Imperial standard weight, but it is still optional with the physician in *prescribing* to use the symbols ℥ (scruple) and ℥ (drachm), the former representing 20 and the latter 60 grains. In the measurement of liquids the Imperial measure is used for higher denominations, and the fluid ounce and its subdivisions into fluid drachms and minims for the lower denominations of volume.'

MEASURES

The Imperial gallon contains 277 274 cubic inches of distilled water at 60° F

℥ 1 gallon	= 8 pint,	weighing 10 pounds,	contains 76,800 minims
℥ 1 pint	= 20 fluid ounces	1½ "	" 9,600 "
℥ oz 1 fluid ounce	= 8 fluid drachms	437 5 grains	" 480 "
℥ dr 1 fluid drachm	= 60 minims	54 68 "	" 60 "
℥ m 1 minim		91 grain	" 1 minim

It must be remembered that the minim is less than the grain measure, 109 7148 minims (taken as 110 minims throughout *B.P.* '94) = the volume of 100 grains of Water at 60° F (15 5° C).

To find the capacity in gallons of any rectangular vessel, multiply the length in inches by the breadth, and the product by the depth in inches, then divide the total by 277 274, which is the number of cubic inches contained in the gallon.

To find the capacity in gallons of a cylindrical vessel, multiply the square of half the diameter in inches by 8 1416 and the resulting figure by the depth in inches, divide the result by 277 274.

Graduated measures may be checked with good weights and scales, and distilled water. Every fluid ounce of distilled water at 60° F (15 5° C) weighs an ounce avoirdupois, but there are two lines on the surface of a liquid; the upper one is that of capillary attraction to the sides of the vessel, the lower one the exact surface of the fluid. This should be on a line with the eye to measure accurately.

The Continental Pharmacopœias give the formulas in parts by weight; in some instances the gramme is indicated as the unit. The formulas in the United States Pharmacopœia are given in grammes and cubic centimetres.

The British Pharmacopœia still gives the formulas in weights and measures, both by the Imperial and the metric systems. Liquids are as a rule ordered by measure, but there is no uniformity in this: for instance, in *Tinctura Terebinthina Aceticum* the Chloral Acetic Acid is weighed, but in *Acetum Cantharidis* it is measured, in *Oxymel* the clarified Honey is weighed, but in *Oxymel Scillæ* it is measured. Glycerin and other fluids, in some preparations are weighed, in others they are measured.

METRICAL MEASURES.

ALSO RELATION OF THE METRICAL MEASURES TO THE
MEASURES OF THE BRITISH PHARMACOPEIA.

LENGTH.

1 Millimetre	the thousandth part of a metre, or 0.001 metre	= 0.03937 inch.
1 Centimetre	the hundredth " "	0.01 " 0.39371 "
1 Decimetre	the tenth part " "	0.1 " 3.93708 inches.
1 Metre	89.87079 inches, or 1 yard 3.37 inches nearly.	

1 Line	$\frac{1}{12}$ inch	
1 Inch	$\frac{1}{12}$ foot	0.02540 metre, or 25.40 millimetres.
12 Inches	1 foot	0.30480 metre, or 30.48 centimetres
36 "	3 feet 1 yard	0.91440 metre.

Length of pendulum vibrating seconds of mean time in the
latitude of London, in a vacuum at the level of the sea } 39.1393 inches.

It is remarkable that the English and French standards, taken from such
different sources, should so nearly agree

The English, from the length of a pendulum vibrating seconds of
mean time, from which the yard (36 inches) is computed } 39.1393.
The French being the ten-millionth part of a quarter of the
earth's meridian, and called a metre } 39.37079

CAPACITY.

1 Millilitre	= 1.00016 cubic centimetre, or the volume of 1 gramme of water at 4°C.	
1 Centilitre	=	10 grammes " "
1 Decilitre	=	" 100 " " "
1 Litre or 1000 Millilitres	} 1.00016 cubic decimetre	" 1000 " (1 kilo) "

1 Cubic Centimetre = .00084 millilitre.

1 Cubic Decimetre (1000 c c) = .00084 litre.

1 Cubic Centimetre = 15.48286 grain measures of 16.9 minims nearly.

1 Litre = 1.7598 pints, 35½ fl. oz. and 11 minims, or 16482.848 grain-measure.

1 Gallon = 4.5460821 litres.

1 Pint = 568.2454 litre, or 568.336 cubic centimetres nearly.

1 Fluid Ounce = .0284128 litre, or 28.417 cubic centimetres nearly

1 Fluid Dram = .008652 litre, or 8.652 cubic centimetres nearly

1 Minim = .000059 litre, or .056 cubic centimetre nearly.

EQUIVALENTS OF ENGLISH WEIGHTS TO FRENCH GRAMMES.

1 pound avoirdupois	7000 grains.	or 16 ounces.	= 458 592 French grammes
	5562 5	or 15	= 425 2425 " "
	5125	or 14	= 396 8925 " "
	5687 5	or 13	= 368 5435 " "
	5250	or 12	= 340 1935 " "
	4812 5	or 11	= 311 8415 " "
	4375	or 10	= 283 495 " "
	3937 5	or 9	= 255 1455 " "
	3500	or 8	= 226 796 " "
	3062 5	or 7	= 198 4465 " "
	2625	or 6	= 170 097 " "
	2187 5	or 5	= 141 7475 " "
	1750	or 4	= 113 398 " "
	1312 5	or 3	= 85 0485 " "
	875	or 2	= 56 699 " "
1 ounce,	437 5	or 1	= 28 3495 " "
	218 75	or $\frac{1}{2}$	= 14 17475 " "
	109 37	or $\frac{1}{4}$	= 7 08737 " "
	15 43		= 1
	1 543		= 1, a decigramme
1 grain,	1		= 0048
	.15 or $\frac{1}{4}$ nearly		= 01, a centigramme
	015 or $\frac{1}{64}$ nearly		= 001, a milligramme (nearly).

MEASURES, EQUIVALENTS OF FRENCH GRAMMES TO
ENGLISH WEIGHTS

1 Litre	= 1 kilogramme, 1000 French grammes	= 35 ounces and 120 grains
	900	= 31 . and 326 $\frac{1}{2}$ "
	800	= 28 . and 96 "
	700	= 24 . and 302 $\frac{1}{2}$ "
	600	= 21 . . and 72 "
	500	= 17 . and 278 $\frac{1}{2}$ "
	400	= 14 . and 48 "
	300	= 10 . and 254 $\frac{1}{2}$ "
	200	= 7 . and 24 "
1 Decilitre = 1 hectogramme,	100	= 3 . and 280 $\frac{1}{2}$ "
	90	= 3 . and 76 $\frac{1}{2}$ "
	80	= 2 . and 359 $\frac{1}{2}$ "
	70	= 2 . and 205 $\frac{1}{2}$ "
	60	= 2 . and 51 "
	50	= 1 . and 334 "
	40	= 1 . and 179 $\frac{1}{2}$ "
	30	= 1 . and 25 $\frac{1}{2}$ "
	20	= . . 808 $\frac{1}{2}$ "
1 Centilitre = 1 decagramme	10	= . . 154 $\frac{1}{2}$ "
	5	= . . 77 $\frac{1}{2}$ "
1 Millilitre = *1 gramme,	1	= . . nearly 15 $\frac{1}{2}$ "
	5	= . . 7 $\frac{1}{2}$ "
1 decigramme,	.1	= . . 1 $\frac{1}{2}$ "
	.05	= . . 1 $\frac{1}{2}$ "
1 centigramme,	01	= . . 1 $\frac{1}{2}$ "
	005	= . . 1 $\frac{1}{2}$ "
1 milligramme,	001	= . . 1 $\frac{1}{2}$ "

* A *Millilitre* is the volume of one gramme of Distilled Water at its greatest density, 4° C (39 2° F). A *Cubic Centimetre* is the volume of the same weight of water at 60° F (15 5° C).

ALCOHOL TABLE.

Specific Gravity at 60° F (15.6° C)	Absolute Alcohol by weight Per cent	Absolute Alcohol by volume Per cent	Specific Gravity at 60° F (15.6° C)	Absolute Alcohol by weight Per cent	Absolute Alcohol by volume Per cent
1.000	0.00	0.00	.891	60.67	68.33
.998	1.06	1.34	.892	61.50	69.11
.996	2.28	2.86	.890	62.36	69.92
.994	3.41	4.27	.888	63.26	70.77
.992	4.62	5.78	.886	64.13	71.58
.990	5.87	7.32	.884	65.00	72.38
.988	7.21	9.01	.882	65.83	73.15
.986	8.61	10.73	.880	66.50	73.93
.984	10.08	12.49	.878	67.54	74.70
.982	11.62	14.37	.876	68.38	75.45
.980	13.15	16.24	.874	69.21	76.20
.978	14.82	18.26	.872	70.04	76.94
.976	16.46	20.24	.870	70.84	77.64
.974	18.08	22.18	.868	71.67	78.36
.972	19.67	24.08	.866	72.50	79.12
.970	21.31	26.01	.864	73.30	79.86
.968	22.83	27.86	.862	74.13	80.60
.966	24.38	29.67	.860	75.11	81.40
.964	25.86	31.40	.858	76.04	82.19
.962	27.21	32.98	.856	76.88	82.90
.960	28.56	34.51	.854	77.71	83.60
.958	29.87	36.04	.852	78.52	84.27
.956	31.00	37.34	.850	79.32	84.93
.954	32.25	38.75	.848	80.13	85.59
.952	33.47	40.14	.846	80.96	86.23
.950	34.52	41.32	.844	81.76	86.93
.948	35.50	42.40	.842	82.54	87.55
.946	36.56	43.56	.840	83.31	88.16
.944	37.67	44.70	.838	84.08	88.76
.942	38.78	46.02	.836	84.88	89.38
.940	39.80	47.13	.834	85.65	89.99
.938	40.80	48.21	.832	86.42	90.58
.936	41.80	49.29	.830	87.19	91.17
.934	42.76	50.31	.828	87.96	91.75
.932	43.71	51.32	.826	88.76	92.36
.930	44.64	52.29	.824	89.54	92.94
.928	45.55	53.24	.822	90.29	93.49
.926	46.46	54.19	.820	91.00	94.00
.924	47.36	55.13	.818	91.71	94.51
.922	48.27	56.07	.816	92.44	95.03
.920	49.16	56.98	.814	93.18	95.55
.918	50.09	57.92	.812	93.90	96.08
.916	50.96	58.80	.810	94.62	96.55
.914	51.79	59.68	.808	95.32	97.02
.912	52.68	60.52	.806	96.04	97.51
.910	53.57	61.40	.804	96.70	97.94
.908	54.48	62.31	.802	97.37	98.37
.906	55.41	63.24	.800	98.03	98.80
.904	56.32	64.14	.798	98.66	99.16
.902	57.21	65.01	.796	99.29	99.55
.900	58.05	65.81	.794	99.94	99.96
.898	58.95	66.69	.7938	100.00	100.00
.896	59.83	67.53			

BEAUME'S HYDROMETER COMPARED WITH THE SPECIFIC GRAVITY OF LIQUIDS HEAVIER THAN WATER

1 000 BEING TAKEN AS THE SPECIFIC GRAVITY OF DISTILLED WATER AT
15.57 CENTIGRADE = (60) FAHRENHUIT

Beaume	Sp. Gr.	Beaume	Sp. Gr.
0	1 000	39	1 967.
1	1 007	40	1 980
2	1 014	41	1 991
3	1 021	42	1 407
4	1 028	43	1 421
5	1 036	44	1 435
6	1 043	45	1 449
7	1 051	46	1 464
8	1 058	47	1 479
9	1 066	48	1 494
10	1 074	49	1 510
11	1 082	50	1 526
12	1 090	51	1 542
13	1 098	52	1 558
14	1 107	53	1 575
15	1 115	54	1 593
16	1 124	55	1 610
17	1 133	56	1 628
18	1 142	57	1 647
19	1 151	58	1 666
20	1 160	59	1 685
21	1 169	60	1 705
22	1 179	61	1 725
23	1 188	62	1 746
24	1 198	63	1 767
25	1 208	64	1 789
26	1 218	65	1 811
27	1 229	66	1 834
28	1 239	67	1 857
29	1 250	68	1 882
30	1 261	69	1 906
31	1 272	70	1 932
32	1 283	71	1 958
33	1 294	72	1 984
34	1 306	73	2 011
35	1 318	74	2 040
36	1 330	75	2 069
37	1 342	76	2 099
38	1 355	77	2 180

SPECIFIC GRAVITY of Syrups, etc., may be tested with a ten ounce measure
Ten measured ounces of simple syrup should weigh nearly thirteen ounces and
146 grains representing the sp. gr. 1 330.

MATERIA MEDICA TABLE

See Notes

R. P. Name	Obtained from	Natural Order	Geographical Source
Acacia Cortex.	Acacia Arabica and Acacia decurrens	Leguminosæ (Fabaceæ)	Africa, Australia and India
Acacia Gummi	Acacia Senegal, and other species	"	Rhodesia in East of Africa
Acalypha	Acalypha Indica	Euphorbiaceæ	India
Aconiti Folia	Aconitum 'Napellus'	Ranunculaceæ	Britain
Aconiti Radix	"	"	"
Aconitina	"	"	"
Adeps Iamæ	Ovis Ariæ	Ungulata	Domesticated everywhere
Adops	Sus Scrofa	"	"
Adhatoda	Adhatoda Verrea (Justicia Adhatoda)	Acanthaceæ	India
Agropyrum	Agropyrum repens	Graminaceæ	Europe, North America, and Asia
Ajowan Oleum	Carum Copticum	Umbellifera (Apiaceæ)	India, Egypt and Persia
Aloe Barbadosensis	Aloe Chinensis, A vera, and probably other species	Liliaceæ	(Dutch West Indian Islands) and Barbados
Aloe Socotrina	Aloe Perryi and probably other species	"	Socotria (shipped by way of Bombay and Zanzibar)
Alstonia	Alstonia scholaris Alstonia constricta	Apocynaceæ	India, the Philippine Islands, Australia
Ammoniacum	Dorema Ammoniacum and other species	Umbellifera	Persia
Amygdala Amara	Prunus Amygdalus (var amara)	Rosaceæ	(South of France, Sicily, and Morocco)
Amygdala Dulcis	Prunus Amygdalus (var dulcis)	"	Spain, Portugal, and South of France
Amygdala Oleum	Both of the above	"	" " "
Amylum	Triticum sativum (wheat). Zea 'Mays' (maize). Oryza sativa (rice)	Graminaceæ	(Cultivated in various parts of the world)
Andrographis	Andrographis paniculata	Acanthaceæ	India, Ceylon, Java
Anethi Fructus	Poucedanum graveolens	Umbellifera	(Ireland, Middle and South of Europe)
Anisi Fructus	Pimpinella Anisum.	"	(Central and Southern Europe, etc.)
Anisi Oleum	Illicium anisatum	Magnoliaceæ	(Distilled in Europe and in China)
Anthemidis Flores	Anthemis nobilis	Compositæ (Asteraceæ)	(Britain and Belgium, cultivated)
Anthemidis Oleum	" "	"	(Britain and Belgium, and France, etc.)
Apomorphina Hydrochloridum	Morphina or Codeina	see Opium	see Opium
Arachidis Oleum	Arachis hypogæa	Leguminosæ	(Tropical Africa, China, India, etc., cultivated)

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Parts used

Preparations into which it enters

Dried bark	Decoctum Acacie Corlicis
Gum exuded from stem and branches	Mucilago Acacie Pulv Amygdalæ Co and Pulv Tragaanth Co All Trochisci
Fresh and dried herb	Extractum Acalyphæ Liquidum, and Succus Acalyphæ
Fresh leaves and flowering tops	
Dried root	Lanimum, and Tinctura Aconiti, and Aconitina
Dried root	Unguentum Aconitine
The purified cholesterol fat of sheep's wool	Adeps Laniæ Hydrosus
Purified fat of the abdomen	Adeps Benzoeatus, Empl Canthar, Pilula Phosphori, Unguenta Aconitine, Atropinæ, Cocainæ, Hydrargyri, Hydrag Nitratis, Iodi, Resinæ, and Veratrinæ
Fresh and dried leaves	Extractum Adhatodæ Liquidum, Succus Adhatodæ, and Tinctura Adhatodæ
Dried rhizome	Decoctum Agropyri, and Extractum Agropyri Liquidum
Distilled from fruit	
Juice of the leaf evaporated to dryness	Mom Decoct Aloe Co Ext Aloe Barb, Ext Coloc Co, Pil Aloe Barb, Pil Aloes et Ferri, Pil Cambogiæ Comp, Pil Colocynthidis Co, Pil Colocynthidis et Hyoscyami Tinctura Aloes
Juice of the leaf evaporated to dryness	Mom Ext Aloes Socotrine, Pil Aloes et Asafetidæ, Pil Aloes et Myrrha, Pil Aloes Socot, Pil Rhei Co, Tinct Benzoin Co
Dried bark	Infusum Alstoniæ, and Tinctura Alstoniæ
Gum resinous exudation from the flowering and fruiting stem	Emplastrum Ammoniaci c Hydrargyro, Emplast Galbani, Mistura Ammoniaci, Pilula Scillæ Co, Pil Ipecac c Scillâ.
Ripeseed	Oleum Amygdalæ
Ripeseed	Oleum Amygdalæ, Pulvis Amygdalæ Compositus, Mistura Amygdalæ
Expressed oil from seeds	Ol Phosphoratum, Unguenta Aquæ Rosæ, and Cetacei
Starch	Glycerinum Amyli, Pulvis Tragacanth Co
Dried plant	Infusum Andrographidis, Laquor Andrographidis Concentratius, and Tinctura Andrographidis.
Dried ripe fruit	Aqua Anethi, Oleum Anethi
Dried ripe fruit	Aqua Anisi, Oleum Anisi
Oil distilled from fruits	Spiritus Anisi, Tinct Camph Co, Tinct Opi Ammoniacata
Dried expanded flower heads or capitula	Extractum and Oleum Anthemidis The latter is also used in the preparation of the Extract
Volatile oil	Extractum Anthemidis
Salt of the alkaloid	Injection Apomorphinæ Hypodermica
Expressed oil, without heat	

R.P. Name	Obtained from	Natural Order	Geographical Source
Araroba	Andia 'Araroba'	Leguminosae	Brazil (Bahia)
Areca	Areca Catechu	Palmaceae	Malayan Archipelago
Aristolochia	Aristolochia Indica	Aristolochiaceae	India and Holland
Arnicae Radix	Cochlearia <i>Arnica</i>	Cruciferae (Brassicaceae)	Britain, cultivated
Arnicae Flores	Arnica montana	Compositae	Mountainous parts of Middle and Southern Europe
Arnicae Rhizoma	" "	" "	" "
Asafoetida	<i>Perula fetida</i> , and " " " "	Umbelliferae	Afghanistan and Persia
Atropina	<i>Belonia</i> <i>Atropa</i> <i>Folia</i>	Solanaceae	Britain, Germany, Austria, etc.
Aurantii Floris Aqua	<i>Citrus Aurantium</i> var 'Bigaradia'	Rutaceae	South of Europe
Aurantii Cortex	" "	" "	" "
Aurantii Cortex Indicus	<i>Citrus Aurantium</i>	" "	India and Ceylon
Asadirachta Indica	<i>Melia Asadirachta</i>	Meliaceae	Southern India, Ceylon
Balsamum Peruvianum	<i>Myroxylon Peronae</i>	Leguminosae	San Salvador, in Central America
Balsamum Toluianum	<i>Myroxylon Toluifera</i>	" "	New Granada
Bebeeru Cortex	<i>Nectandra Rodiei</i>	Lauraceae	Guiana
Belle Fructus	<i>Aegle Marmelos</i>	Rutaceae	India
Belladonna Folia	<i>Atropa Belladonna</i>	Solanaceae	Britain
Belladonnae Radix	" "	" "	Britain, Germany, etc.
Benzoinum	<i>Styrax Benzoin</i> and other species	Styracaceae	Siam and Sumatra
Berberidis Cortex	<i>Berberis vulgaris</i>	Berberidaceae	Britain
Berberis	<i>Berberis aristata</i>	" "	India and Ceylon
Betel	<i>Piper Betle</i>	Piperaceae	India, Ceylon, etc.
Bryoniae Radix	<i>Bryonia dioica</i> <i>Bryonia alba</i>	Cucurbitaceae	England and Central and Southern Europe
Buchu Folia	<i>Barosma betulina</i>	Rutaceae	Cape Colony
Butea Gummi	<i>Butea frondosa</i>	Leguminosae	India
Butea Seminum	" "	" "	" "
Cardianum Oleum	<i>Juniperus Oxycedrus</i> and other species	Coniferae (Pinaceae)	Southern Europe
Caffeina	<i>Camellia Thea</i> <i>Coffea Arabica</i>	Camelliaceae Rubiaceae	China, Japan, and Upper India Cultivated throughout the Tropics (native of Abyssinia)

Parts used	Preparations into which it enters
{ A substance found in cavities in the trunk of Andira Araroba, freed as much as possible from fragments of wood, dried and powdered }	Chrysaroban, of Chry-aroban, Ung Chrysaroban
Seeds	
Dried stem and root	Liquor Aristolochie Concentratus, and Tinctura Aristolochie
Fresh root	Spiritus Armoracis Compositus
Dried flower-heads	Tinctura Arnicæ Florum
Dried rhizome and roots	Tinctura Arnicæ
Gum resin	{ Pil Aloes et Asafetide, Pil Galbani Co, Spiritus Ammoniacæ Fœtidus, Tinctura Asafetide
Alkaloid	Unguentum Atropine, Atropine Sulphas, Lamelle Atropine and Liquor Atropine Sulphatis
Distilled water of flowers	{ Syrupus Auranti Floris, Mistura Olei Ricini, Syrupus Calcii Lactophosphatis
Dried outer part of the rind	{ Fresh Peel Tinctura Auranti, Vinum Aurant, Syrupus Auranti, Tinct Quina, Syrupus Aromaticus, and Syrup Cascara Vinat Dried Peel Inf Auranti, Inf Aurantil Comp, Inf Gentian Comp, Tinct Cinchon Comp, Tinct Gentian Comp, Spirit Armoracis Comp
{ The fresh and dried outer part of the rind }	
Dried bark of the stem	Infusum Azadirachtæ Indice, and Tinctura Azadirachtæ
Balsam, from the trunk	
Balsam, from the trunk	{ Mist Ammoniaci, Syrupus Tolutanus, Tinct Tolutana, Tinct Benzoin Co, Tolu Basis, Troch Acidi Carbolici, Troch Morphine, Troch Morphine et Ipecacuanhæ
Dried bark	
Fresh half-ripe fruit	Extractum Belladonnæ Liquidum
Fresh leaves and branches	Extract Belladonnæ Viride, Succus Belladonnæ, Atropina, Atropina, Emplast Belladonnæ, Extract Belladonnæ Alcoholium, Extractum Belladonnæ Liquidum, Linimentum Belladonna Tinct Belladonna, Ung Belladonnæ, Suppos Belladonnæ
Dried root	{ Acidum Benzoicum, Adeps Benzoeatus (and Ointments containing it), Tinct Benzoin Co, Ung Cæteri Extractum Berberidis Fluidum, Berberine Phosphas Liquor Berberidis Concentratus, and Tinctura Berberidis.
Balsamic resin	
Dried bark	
Dried stem	
Leaves	
Fresh and dried root	Tinctura Bryoniæ
Dried leaves	Infusum Buchu, Tinctura Buchu
Impassated juice from stem	
Seeds	Pulveris Butei Seminum
{ An empyreumatic oily liquid obtained by destructive distillation of the branches and wood }	
{ Alkaloid from leaves of tea or seeds of coffee }	Cassanea Citras, Cassanea Citras Effervescens.

B.P. Name.	Obtained from	Natural Order	Geographical Source
Cajuputi Oleum	Melaleuca leucadendron	Myrtaceæ	{ Imported from Batavia and } Singapore
Calendula Flores	Calendula officinalis	Compositæ	Texas and Southern Europe
Calotropis . . .	{ Calotropis pinnatifida } { Calotropis gigantea }	Asclepiadaceæ	India
Calumbæ Radix	Jateorhiza 'Calumbæ'	Menispermaceæ	{ Eastern Africa, between India } and the Zambesi
Cambogia	Garcinia Hanburyi	Clusiaceæ	Siam
Cambogia Indica	Garcinia Morella	"	India
Camphora . . .	Cinnamomum Camphora	Lauraceæ	{ China (Formosa) and Japan } { (produced in Britain and } elsewhere)
Canellæ Cortex	Canella alba	Canellaceæ	West Indies
Cannabis Indica	Cannabis sativa	Urticaceæ	India
Cantharis . . .	Cantharis vesicatoria	Coleoptera	Spain, France, Sicily, Hun }
Caoutchouc . .	Hevea Brasiliensis	Euphorbiaceæ	{ ary, and Southern Russia } { Brazil (Para)
Capsici Fructus .	Capsicum minimum	Solanaceæ	Zanzibar, Sierra Leone, etc.
Carbo Ligni . .	Wood	Various	Britain
Cardamomi Semina	Elettaria Cardamomum	Scitamineæ	India and Ceylon
Carrageen . . .	Chondrus crispus	{ Algae } { (Gelatinifera) }	{ (North Cape to Gibraltar) } { (Eastern Coasts of N. America)
Carui Fructus.	Carum Carvi . . .	Umbellifera	Europe
Carui Oleum . .	" "	"	Central and Northern Europe
Caryophylli Oleum	Eugenia caryophyllata	Myrtaceæ	{ Molucca Islands, Zanzibar, } { and Pemba }
Caryophyllum	" "	"	{ Zanzibar, Penang, Bon } { eolien, etc }
Cascara Sagrada	Rhamnus Purshiana	Rhamnaceæ	California
Cascarilla . . .	Croton Elateria	Euphorbiaceæ	Bahama Islands
Cassia Pulpa	Cassia fistula . . .	Leguminosæ	East and West Indies
Castoreum	Castor Fiber . . .	Rodentia . . .	Siberia and Canada
Catechu	Uncaria 'Gambier'	Rubiaceæ . . .	{ Singapore, and other places } { in the Eastern Archipelago }
Catechu Nigrum } (Pegu Catechu } and Gutch) . . }	Acacia Catechu . . .	Leguminosæ . .	India and Burma
Cera Alba . . .	Apis mellifica . . .	Hymenoptera .	Britain
Cera Flava . . .	" "	"	Britain, etc.
Cetaceum	Physoter macrocephalus	Cetacea	Pacific and Indian Oceans .
Cetraria	Cetraria Islandica	{ Discomycetes } { or Disco- } lichones . . . }	North of Europe

Parts used	Preparations into which it enters
{ Volatile oil distilled from the leaves	{ Spiritus Cajuputi, Lin Crotonis
Dried florlets	
Dried root bark	
{ Dried transversely cut slices of the root	{ Infusum, Liquor Columbe Concentratus, and Tinct Columbe
Gum resin	
Gum resin	Pilula Cambogire Composita
{ A white crystalline substance obtained from the wood	{ Aqua Camph, Linimenta Acroni, Belladonnae, Camph, Camph Ammoniatum, Chloroformi, Hydrargyri, Opii, Saponis, Sinapis, Terbinthinae and Terob Aceticum, Spiritus Camph, Tinct Camph Co, Ung Hydrarg Co
Dried bark	
{ Dried flowering or fruiting top of female plants	{ Extractum Cannabis Ind, Tinctura Cannabis Ind, Tinct Chlorof et Morphinae Comp
The dried beetle	
Prepared milk juice	{ Acetum, Emplastrum, Tinctura and Unguentum Cantharidis, Colloidium Vesicans, Liquor Epispast, Emplast Calofaciens, Liquor Crotonis, Charta Sinapis
Dried ripe fruit	{ Tinctura Capsici, Tinct Chlorof et Morphinae Comp, Ung Capsici
{ Carbonaceous residue from wood	{ Tinct Cardui Co, Ext Coloc Co, Pulv Cinnam Co, Pulv Ciste Arom, Tinct Gentian Co, Tinct Rhei Co, Decoct Aloes Comp, Mist Sennae Comp
The dried ripe seeds	
Dried seaweed	Saccharum, and Gelatina Carrageen
Dried fruit	{ Aqua and Oleum Carui Conf Piperis, Pulvis Opi Compositus, Tinct Cardui Co, Tinct Sennae Co, Pil Aloes Barb
Volatile oil	Pilula Aloes Barbadosensis
Volatile oil	{ Pilula Colocynthis, Composita, Pilula Colocynthis et Hyoscyami
Dried flower buds	{ Infusum and Oleum Caryophylli, Inf Aurant Co, Pulv, Creta Aromat, Pil Coloc Co, Pil. Coloc et Hyoscyami.
Dried bark	{ Ext Cascar Sagrada, Ext Cascar Sagrad Liquid, Syrup Cascar Aromaticus
Dried bark	{ Infusum and Tinctura Cascarilla
Pulp from the pods	
{ Dried preputial follicles and secretion	{ Confectio Sennae
{ An extract of the leaves and young shoots	
Extract from the wood	{ Tinctura Castorei
Honeycomb, wax, bleached	{ Pulvis Catechu Compositus, Tinctura Catechu, Trochiscus Catechu
Honeycomb	{ Pilula Phosphori, Suppositoria Acidi Carbolic, Unguentum Aqua Rosae, and Unguentum Cetacei
{ A concrete fatty substance, mixed with oil, obtained from the head of the sperm whale, purified	{ Emplast Calofaciens, Cantharidis, Pielis; Unguenta Hydrarg Co, Menthol, Pielis Liquid, Rosinum, and Staphisagriae Cera Alba
Dried lichen	
	{ Pilula Phosphori, Suppos Acidi Carbolic, Ung Aquae Rosae, Ung Capsici
	Unguentum Cetacei, Ung Aquae Rosae and Ung Capsici
	Decoctum Cetrariae, Saccharum Cetrariae, Gelatina Cetrariae

R P Name	Obtained from	Natural Order	Geographical Source
Chirata	Swertia 'Chirata' . .	Gentianaceæ	Northern India . . .
Chrysarobinum	See Arnoba.		
Cimicifuga	Cimicifuga racemosa . .	Ranunculaceæ	Canada and U. S. . .
Rhizoma			
Cinchona Flax.	Cinchona Calisaya . .	Rubiacæ	Peru, Southern Peru, and India
Cortex			
Cinchona lancifolia Cortex	Cinchona lancifolia . .		New Granada
Cinchona Rubra Cortex	Cinchona saccharata . .		{ South America, cultivated in East India, Ceylon, Java, etc
Cinnamomi Cortex	Cinnamomum Zeylanicum . .	Lauracæ	Ceylon
Cinnamomi Oleum	"		
Cissampelos	Cissampelos 'Pereira' . .	Menispermaceæ	{ India, East and West Indies, and South Central America . . .
Cocœ Folia	Erythroxylon 'Cocœ' . .	Lanacæ	Peru, Java, and Bolivia . .
Cocœ Hydrochloridum	"	"	"
Cocœus	Cocœus Cacti	Thymelææ	Mexico and Teneriffe . .
Colchici Cornus	Colchicum autumnale . .	Liliacæ	Indy non
Colchici Semina	"	"	"
Colocynthis Pulpa	Citrullus Colocynthis . .	Cucurbitaceæ	{ Northern Africa, Syria, and Spain . . .
Condurango	Gonolobus 'Condurango' . .	Asclepiadaceæ	Equador
Conii Folia	Conium maculatum . .	Umbelliferae	Britain
Conii Fructus	"	"	"
Copaiba	Copaifera Tansdorfii and other species . .	Leguminosæ	Valley of the Amazon, etc . .
Copaibæ Oleum	"	"	"
Coriandri Fructus	Coriandrum sativum . .	Umbelliferae	Germany and Britain, etc . .
Coriandri Oleum			
Cocœinium	Cocœinium fenestratum . .	Menispermaceæ	Ceylon and Southern India
Coto	{ Botanical source uncertain . .	Lauracæ	New Granada
Orocus	Crocus sativus	Iridacæ	Spain and Persia, etc . .
Crotonis Oleum	Croton Tigillum	Euphorbiacæ	{ Hindostan, Ceylon, and Indian Archipelago . .
Jubææ Fructus	Piper Cubeba	Piperacæ	Java
Cucurbitæ Semina Preparata	Cucurbita maxima (Cucurbita Pepo) . .	Cucurbitaceæ	{ Italy, West Indies, Burma, and South America, cultivated . . .
Cuxara	Strychnos toxifera, etc. . .	Loganiacæ	South America
Cuspariæ Cortex	Cusparia febrifuga	Rutacæ	Tropical South America
Cuxa	Brayera anthelmintica . .	Rosacæ	Abyssinia
Cydonium	Pyrus Cydonia	"	Western Asia and Europe . .
Damiana	{ Turnera aphrodisiaca, etc	Turneracæ	Mexico and California . .

Parts used	Preparations into which it enters
Dried plant	Infusum, Liquor Chiratae Concentratus, and Tinctura Chiratae
Dried rhizome and roots	Ext Cinchifugae Liquid, Tinct Cinchifugae
Dried bark	Used in the preparation of the official Salts of Quinine
Dried bark	{ Its use is permitted for the manufacture of the official Salts of Quinine
{ Dried bark of stem and branches of cultivated plants }	{ Ext Cinchon Liquid, Inf Cinchon Acid; Tinct Cinchonae, Tinct Cinchon Comp, Quinine
{ The dried inner bark of shoots from the truncated stocks }	{ Aqua, Oleum, Pulvis Co, and Tinct Cinnamomi, Pulvis Catechu Comp Tinct Catechu, Decoct Hamatoxyli, Pulv Crotae Arom, Pulv Kino Co, Tinct Cardam Co., Tinct Lavand Co, of the Water, Mist Crotae, Mist Guaiaci, Mist Olei Ricini, Mist Spiritus Vin Gallici, Syrupus Aromaticus and Syrupus Cascariae Aromaticus
Volatile oil from bark	{ Spiritus Cinnamomi, of the Compound Powder, Pil. Aloes et Ferri, and Pil Cambogiae Composita, of the Spirit, Acidum Sulphuricum Aromaticum
Dried root	Decoctum Cassiaepae, and Extractum Cassiaepae Liquidum
Dried leaves	Ext Cocae Liquid, Cocaine Hydrochloridum, Cocaine
Salt of the alkaloid	Lamellae Cocaine and Injectio Cocaine Hypodermica
Dried fecundated female insect	Tinctura Cocci, Tinct Cardam Co, Tinct Cinchonae Co
Fresh corn dried	Extractum and Vinum Colchici
Dried ripe seeds	Tinctura Colchici Scammum
{ Dried pulp of the fruit freed from seeds }	{ Extractum Coloc Co, Pil Coloc Co, Pil Coloc et Hyoscyam
Dried bark	
{ Fresh leaves and young branches }	Succus Conii, Unguentum Conii (from Succus)
Dried full-grown unripe fruits	Tinctura Conii
Oleo resin from the trunk	Oleum Copaibae
Distilled oil from oleo resin.	
Dried ripe fruit	{ Oleum Coriandri, Conf Sennae, Syrupus Rhei et Tinct Rhei Co, Syrupus Sennae et Tinct Sennae Co
Volatile oil.	
Dried stem	{ Infusum Coccini, Liquor Coccini Concentratus, and Tinctura Coccini
Bark	
{ The dried stigmas and tops of the styles }	Tinctura Croci, Decoct Aloes Co, Tinct Cinchon Co
Expressed oil from the seeds	Linimentum Crotonis
Dried full grown unripe fruits.	Oleum Cubebae, Tinctura Cubebae
Fresh ripe seeds.	
Extract from plant	
Dried bark	
{ Dried panicles of pistillate flowers. }	Infusum Cuspariae, and Liquor Cuspariae Concentratus
Seeds.	
Leaves	

B.P. Name	Obtained from	Natural Order	Geographical Source
Datura Folia .	<i>Datura fastuosa</i> var. <i>alba</i> . Datura Metel .	Solanaceæ .	India .
Datura	<i>Datura fastuosa</i> var. <i>alba</i> .	" .	" .
Digitalis . . .	<i>Digitalis purpurea</i> .	Scrophulariaceæ .	Britain .
Elettarium . .	<i>Elephantium Elettarium</i> .	Cucurbitaceæ .	Britain (cultivated), Malta .
Elomi	<i>Cassia communis</i> and other species .	Banaceæ .	India, Brazil, and Mexico .
Embelia	<i>Embelia ribes</i> , <i>Embelia</i> <i>robusta</i> .	Myrsinaceæ .	India and East Indies .
Ergota	<i>Secale cereale</i> .	Graminaceæ .	Europe and the Canaries .
Erythrophloeum .	<i>Erythrophloeum Gume</i> <i>ense</i> .	Leguminosæ .	Western Africa
Eucalypti Oleum	<i>Eucalyptus globulus</i> , and other species .	Myrtaceæ .	Australia, etc .
Eucalypti Gummum	<i>Eucalyptus rostrata</i> and other species .	" .	"
Euonymi Cortex .	<i>Euonymus alatus</i> <i>pureus</i> .	Celastraceæ .	United States .
Euphorbia pilulifera	<i>Euphorbia pilulifera</i> .	Euphorbiaceæ .	(Queensland and Tropical) America
Euphorbium . .	<i>Euphorbia resinsifera</i> .	" .	Morocco
Fel Bovinum Purificatum	<i>Bos Taurus</i> .	Ungulata .	Domesticated everywhere .
Ficus	<i>Ficus Carica</i> .	Urticaceæ .	Smirna
Felix-Mas . . .	<i>Aspidium Felix-mas</i> .	Filices . .	Britain
Feniculi Fructus	<i>Fenniculum capillare</i> .	Umbellifera .	(Central and Southern) Europe, also India and Japan, etc.
Fucus	<i>Fucus vesiculosus</i> .	Alga (Fucaceæ) .	Britain
Galbanum . . .	<i>Ferula galbaniflua</i> and probably other species .	Umbellifera .	Persia
Galla	<i>Quercus infectoria</i> .	Umbellifera (Corylaceæ) .	Asia Minor .
Gaultheria Oleum	<i>Gaultheria procumbens</i> (in wax) <i>Betula lutea</i> (bark) .	Ericaceæ Botulaceæ .	United States and Canada .
Gelsemium Radix	<i>Gelsemium nitidum</i> .	Loganiaceæ .	Southern part of the United States of America
Gentiana Radix .	<i>Gentiana lutea</i> .	Gentianaceæ .	Central and Southern Eu- rope (mountains)
Glycyrrhiza Radix	<i>Glycyrrhiza glabra</i> and other species .	Leguminosæ .	England, France, Spain, Siberia, Russia, and Persia .
Gossypii Radicis Cortex	<i>Gossypium herbaceum</i> .	Malvacæ .	Imported from the United States
Gossypium . . .	<i>Gossypium Barbadosense</i> and other species .	" .	Warm and Tropical regions .
Graminis Citrati Oleum	<i>Andropogon citratus</i> .	Graminaceæ .	(India, Malay Peninsula, Ceylon, etc. (cultivated) . .
Granati Cortex .	<i>Punica Granatum</i> .	Lythraceæ .	Shores of the Mediterranean and Central Asia .
Grindelia . . .	<i>Grindelia camporum</i> .	Compositæ .	Southern part of the United States of America
Gualaci Lignum .	<i>Gualacum officinale</i> or <i>Gualacum sanctum</i> .	Zygophyllaceæ .	St. Domingo and Jamaica .

Parts used

Preparations into which it enters

Dried leaves

Dried seeds

Dried leaves

Nearly ripe fruit

Oleo resinous exudation

Fluit

The sclerotium of *Claviceps purpurea*, originating in the ovary of *Saccharale*

Bark

Distilled oil from the fresh leaves

A ruby coloured exudation

Dried root bark

Dried herb

Concrete resinous juice

The purified ox bile

Dried fleshy receptacles

Dried rhizome

Dried ripe fruit

Dried seaweed

Gum-resin

Excrecences caused by the punctures and deposition of an egg or eggs of *Cynips Galla tinctoria*

Distilled oil

Dried rhizome and roots

Dried rhizome and roots

The peeled root and peeled subterranean stem

Dried root bark

Hairs of the seed

Distilled oil 'Oil of Verbena'

Dried bark of the stem and root

Dried leaves and flowering tops

Heart wood

Resin from the stem

Tinctura Daturæ Seminum

Infusum and Tinctura Digitalis

Elatium, Pulvis Elatium Compositus

Extractum Ergotæ, Extractum Ergotæ Liquidum, Infusum Ergotæ, and Tinctura Ergotæ Ammoniata, Ergotinum, Inject Ergotæ Hypodermi

Unguentum Eucalypti

Trochiscus Eucalypti Gummi

Extractum Euonymi Siccum

Confectio Sennæ

Extractum Filicis Liquidum

Aqua Fœniculi, Pulv Glycyrrhizæ Co

Pil Galbani Co

Acidum Gallicum and Tannicum, Tinct Gallæ, Ung Gallæ, and Ung Gallæ c Opio

Tinct Gelsemi

Extractum, Infusum Co, and Tinct Gentianæ Co

Extract, Ext Liquid, Liquor Sarsæ Comp Conc, and Pulv Glycyrrh Co, Pil Hydr Of the Extract, Conf Sennæ, and Decoct Aloes Comp Of the Liquid Extract, Mistura Sennæ Comp, and Tinctura Aloes

Decoctum Gossypii Radicis Corticis, and Extractum Radicis Corticis Liquidum

Pyroxylin

Decoctum Granati Corticis

Extractum Grindeliæ Liquidum

Liquor Sarsæ Compositus Concentratus

Mist Guaiaci, Pil Hydrarg Subchlor Co, Tinct Guaiaci Ammon Trochiscus Guaiaci Resinæ

B.P. Name	Obtained from	Natural Order.	Geographical Source
Gummi Indrieum	Anogeissus latifolia	Combretaceae	India and Ceylon.
Gynocardia Oleum	Taraktogenos Kurzii	Bixaceae	India, Eastern
Hæmatoxylum Lig- num	Hæmatoxylum Campe- chianum	Leguminosae	Campechy, Honduras, and Jamaica
Hamamelidis Cor- tex	Hamamelis Virginiana	Hamamelidaceae	United States and Canada.
Hamamelidis Folia	"	"	"
Hemidesmus Radix	Hemidesmus Indicus	Asclepiadaceae	India and Ceylon
Hirudo	Sanguisuga {medicinalis (speckled) offenalis (green)}	Hirudinea	Spain, France, Italy, Hungary
Hirudo Austriaca	Hirudo quadrata (Linn.) (Lamellidella quad- rata)	"	Australia
Hydrastis Rhizoma	Hydrastis Canadensis	Ranunculaceae	United States and Canada.
Hygrophila	Hygrophila spinosa	Acanthaceae	India
Hyoscyami Folia	Hyoscyamus niger	Solanaceae	Britain and Germany, etc
Ignatia amara	Stychnos Ignatii	Loganiaceae	Philippines
Ipecacuanha Radix	Psychotria Ipecacuanha	Rubiaceae	Brazil and Selangor
Iris	Iris versicolor	Iridaceae	United States
Isapgula	Isotriaena ovata	Plantaginaceae	India and Persia
Jaborandi Folia	Piper Jaborandi	Rutaceae	Brazil (Pernambuco)
Jalapa	Ipomoea 'Purga'	Convolvulaceae	Mexico (cultivated in Ja- maica and India)
Juniperi Oleum	Juniperus communis	Coniferae	North of Europe, indigenous
Kaladana	Ipomoea hederacea	Convolvulaceae	India and Persia
Kamala	Mallotus Philippinensis	Euphorbiaceae	Ceylon, the Philippines, etc.
Kava Rhizoma	Piper Methysticum	Piperaceae	China, Australia, etc.
Kino	Pterocarpus marsupium	Leguminosae	Sandwich Islands
Kino Eucalypti	" eucalyptia	Myrtaceae	Malabar
Kola	Cola acuminata	Sterculiaceae	Western Australia
Krameria Radix	Krameria radialis (Peruvian) Krameria argentea (Pará)	Polygalaceae	West Coast of Africa.
Lactuca	Lactuca virosa	Compositae	Peru and Brazil
Larici Cortex	Larix Europaea	Coniferae	Western, Central and South- ern Europe and Britain
Laurocerasus Folia	Prunus Laurocerasus	Rosaceae	England, Southern and Cen- tral Europe.
Lavandula Oleum	Lavandula vera	Labiatae (Lamiaceae)	Britain, cultivated England, cultivated (also the Western shores of the Mediterranean)

Parts used	Preparations into which it enters
{ Gummy exudation from the wood	Mucilago Gummi Indici
Fatty oil expressed from seeds	Unguentum Gynocardiae
Heart wood	Decoctum Hamamelidis
Dried bark	Tinctura Hamamelidis
Leaves, fresh and dried	{ Of the Dried Leaves, Extractum Hamamelidis Liquidum, Unguentum Hamamelidis (from Liquid Extract) Of the Fresh Leaves, Liquor Hamamelidis
Dried root	Syrupus Hemidesmi
Leech (annelid)	
Leech (annelid)	
Dried rhizome and roots	Extractum Hydrastis Liquidum, Tinctura Hydrastis
Dried herb including the root	Decoctum Hygrophilae
{ Fresh leaves and flowers, with branches	Extractum Viride, Succus, and Pil Coloc et Hyoscyam
{ Dried leaves and flowering tops	{ Tinctura Hyoscyami, Hyoscyam Hydrobrom, Hyoscyamine Sulphas
Seeds (dried)	
Dried root	{ Tinctum Ipecac, Pil Ipecac e Scilla, Pulv Ipecac Co, Trochiscus Ipecac, Trochiscus Morphinae et Ipecac, Vinum Ipecac, Extractum Ipecacuanha Liquidum
Rhizome and roots	
Dried seeds	Decoctum Ispaghulae
Dried leaflets	Extract Liquidum, and Tinct Jaborandi; Pilocarpinae Nitras
Dried tubercles	{ Extractum, Pulv Co, Resina, and Tinctura Jalapa, Pilula Scammoni Composita, Pulvis Scammoni Comp
{ Oil from the full grown unripe green fruit	Spiritus Juniperi, and Mistura Crocoti
Dried seeds	{ Pulvis Kaladanae Compositus, Resina Kaladanae, and Tinctura Kaladanae
{ Minute glands and hairs from the fruits	
Rhizome without the roots	Extractum Kava Liquidum
{ Juice from the trunk evaporated to dryness	Pulvis Compositus, Tinct Kino, Pulv Catechu Co
Exudation from the stem	
The seeds	
Dried root	{ Extractum, Infusum, Tinctura Krameriae, Pulv Catechu Co, Liqueur Krameriae Concentratus, Troch Krameriae; Troch Krameriae et Cocumae
{ Fresh herb	
Bark (dried)	
Fresh leaves	Aqua Laurocerasi
Distilled oil from flowers	{ Spiritus and Tinctura Lavandulae Comp, Lin Camph Ammoniatum, Tinct Lavand Comp is contained in Liqueur Arsenicalis

R.P. Name	Obtained from	Natural Order	Geographical Source
Limonis Cortex	<i>Citrus medica</i> var. <i>"Limonium"</i>	Rutaceæ	South of Europe
Limonis Succus	"	"	" and West Indies
Linum	<i>Linum usitatissimum</i>	Linacæ	Britain, Holland, Russia, etc.
Lobelia	<i>Lobelia inflata</i>	Campanulacæ	North America
Lupulinum	<i>Humulus Lupulus</i>	Urticacæ	England, imported from Europe, United States, and Canada
Lupulus	"	"	"
Lycopodium	<i>Lycopodium clavatum</i> and other species	Lycopodiaceæ	Great Britain, Central and Northern Europe, etc.
Manna	<i>Fraxinus Ornus</i>	Oleaceæ	Sicily and Calabria
Mastiche	<i> Pistacia Lentiscus</i>	Anacardiaceæ	The Island of Sicily
Matico	<i>Piper acutifolium</i>	Piperacæ	Peru, etc.
Mel Depuratum	<i>Apis mellifica</i>	Hymenoptera	Universally domesticated
Menthæ piperitæ Oleum	<i>Mentha piperita</i>	Labiatæ	Britain and the United States, etc.
Menthæ viridis Oleum	<i>Mentha viridis</i>	"	Britain and Germany, etc.
Menthol	<i>Mentha arvensis</i> vars. <i>piperascens</i> and <i>glabrata</i> <i>Mentha piperita</i>	"	China and Japan United States
Mezeri Cortex	<i>Daphne</i> { <i>Mazoreum</i> <i>Laurula</i> <i>Gnidium</i> }	Thymelacææ	Mountainous parts of Europe
Morphinæ Acetas	Opium	see Opium	see Opium
Morphinæ Hydrochloridum	"	"	"
Morphinæ Tartras	"	"	"
Morrhue Oleum	<i>Gadus Morhua</i>	Telosteæ	Coasts of Norway, France, and England, Newfoundland and Labrador
Moschus	<i>Moschus moschiferus</i>	Ungulata	Native of Central Asia, imported from China.
Mylabris	<i>Mylabris phalerata</i> and <i>M. Cichorii</i>	Coleoptera	China, Southern Europe, and India
Myristica	<i>Myristica fragrans</i>	Myristicacææ	Banda Islands of the Malay Archipelago and Sumatra, etc.
Myristicæ Oleum	" "	"	" "
Mysobalanum	<i>Terminalia Chebula</i>	Combretacææ	East Indies
Myrrha	<i>Balsamodendron</i> { <i>Myrrha</i> and probably other species }	Burseracææ	Somali-land and Arabia Felix
Nux Vomica	<i>Strachnos Nux vomica</i>	Loganiacææ	India, reported from Java, Ceylon, and Cochinchina

Parts used	Preparations into which it enters
Flesh outer part of the pericarp of the fruit	Oleum, Sympus, Tinctura Jambonis, Inf Aurant Co, Inf Gentian Co The Oil is contained in Lin Pot Iodid, c Saponis, Spir Amm Aromat, Tinct Guaiac Amm and Tinct Valer Amm
Freshly expressed juice of the ripe fruit	Syrupus Jambonis, Acidum Citricum
Dried ripe seeds, entire and reduced to a coarse powder	Linum Contusum, Oleum Jamb
Dried flowering herb	Tinct Lobelia Etherea
Glands from the strobiles	
Dried strobiles	Infusum, Tinctura Lupuli, Lupulinum
The minute yellow spores	
Concrete saccharine exudation	
Concrete resinous exudation	
Dried leaves	
Saccharine secretion in honey comb	Mel Borac, Oxymel, Oxymel Scille, Conf Piper
Oil distilled from fresh flowering peppermint	Aqua, and Spiritus Menthe Piperite, Pil Rhei Co, Tinct Chloroform et Morphine Comp
Oil distilled from fresh flowering spearmint	Aqua Menthe Viridis
(A crystalline substance obtained by cooling the oil)	Emplastrum Menthol
Dried bark	Liquor Sarsa Compositus Concentratus
Salt of the alkaloid	Liquor Morphine Acetatis
Salt of the alkaloid	Liquor Morph Hydrochlor, Suppos Morph, Tinct Chloroform et Morph Comp, Trochiscus Morph, also of Ipecac
	Injectio Morphine Hypodermica, and Liquor Morphine Tartratis
Oil of fresh liver of the cod	
(Dried secretion from the preputial follicles)	
The dried beetle	Acetum Mylabridis, Emplastrum Mylabridis, Emplastrum Calefaciens Mylabridis, Liquor Epispasticus Mylabridis, Unguentum Mylabridis
Dried seed divested of its testa	Oleum, and Oleum Myristice Expressum, Pulv Catechu Co, Pulv Cate Aromat, Sp Armoracis Co, Tinct Lavand Co
Oil distilled from seed	Pil Aloes Sacot, Spir Ammon Arom, Spir Myristice, Tinct Guaiac Amm and Tinct Valer Amm, of the Spirit, Mist Ferri Comp
Dried immature fruits	Unguentum Myrobalani, Unguentum Myrobalani c. Opio.
Gum-resin (from the stem)	Tinct Myrrh, Pil Aloes et Myrrh, Decoct Aloes Co, Mist Ferri Co, Pil Galban Co, Pil Rhei Co
Dried ripe seeds.	Extractum, Extractum Liquidum, and Tinctura Nucis Vomice, Strychnina

B. P. Name	Obtained from	Natural Order	Geographical Source
Olive Oleum .	Olea Europaea	Oleaceae	South of Europe
Oliveri Cortex.	Cinnamomum Oliveri	Lauraceae	{Australia, India, Burma, North America, and Southern United States . }
Opium . .	Papaver somniferum	Papaveraceae	Asia Minor (Smyrna, etc.)
Papaveris Capsulae	"	"	Britain and Asia Minor
Parcelm Radix	{Chondrodendron tomentosum	Menispermaceae	Brazil . .
Pepsinum	{Sus Scrofa Ovis Aries Bos Taurus	Ungulata	Domesticated everywhere .
Physostigmatis Semen	Physostigma venenosum	Leguminosae	Western Africa
Pterorhiza .	Pterorhiza 'Kuriou'	Scrophulariaceae	Northern India .
Pterotoxinum	Anamirta paniculata	Menthastraceae	Eastern India and the Malayan Island . }
Pimenta .	Pimenta officinalis	Myrtaceae	Jamaica
Pimentis Oleum	" "	"	{West Indies, Mexico, and Jamaica, etc. . . . }
Pinii Oleum .	Pinus pumilio	Coniferae	Mountains of Central Europe
Piper Nigrum .	Piper nigrum	Piperaceae	East Indies
Pice Burgundica	Picea excelsa .	Coniferae	Germany
Pice Laguida	{Pinus sylvestris and other species	"	Scotland, Russia, Denmark, and Norway . }
Podophylli Indici Rhizoma	Podophyllum pmodi	Berberidaceae	Indigenous to the Himalayas
Podophylli Rhizoma	Podophyllum peltatum	"	North America
Pruni Virginianae Cortex	Prunus serotina . .	"	" "
Prunum . . .	Prunus domestica .	Rosaceae	South of France .
Pterocarpis Lignum	Pterocarpus santalinus	Leguminosae	Madras and Ceylon . . .
Pyrethri Flores	{Pyrethrum cinerarius- folium, P. roseum and P. carneum	Compositae	{California, Dalmatia, and the Caucasus . . . }
Pyrethri Radix .	Anacyclus Pyrethrum .	"	Algeria
Quassiae Lignum	Picramnia excelsa .	Simarubaceae	Jamaica
Quillaise Cortex	Quillaja 'Saponaria'	Rosaceae	Chile and Peru
Quininae Hydrochlor . . .	{Various species of Cin- chona and Remijna }	Rubiaceae	{South America, India, Ceylon, and Java . . }
Quininae Sulphas .	" "	"	" "
Resina . . .	Various species of Pinus	Coniferae . .	North America
Rhamni Purshiani Cortex	See Oascara Sagrada.		
Rhei Radix .	{Rheum palmatum, Rheum officinale, and probably other species }	Polygonaceae .	China and Tibet .
Rheados Petala .	Papaver Rhoeas . .	Papaveraceae .	Britain

Parts used	Preparations into which it enters
Expressed oil from the ripe fruit	Emplastrum Ammoniac e Hydrag, Hydriac Piceis, Plumbi, Linimenta Ammoniac, Cicutis and Camphorae, Sapo Durus and Molles, Unguentum Capsici, Hydrag, Comp, Hydrag Nitratis, and Resin e
Dried bark	Tinctura Olivæ Corticis
Inspissated juice from unripe capsules	Preparations many Tida Opium
Nearly ripe dried fruits	
Dried root	Extractum Pareiræ Liquidum
An enzyme from the mucous lining of the fresh and healthy stomach of the pig, sheep or calf	Glycorium Pepsinæ
Ripe seeds	Extractum Physostigmatis, Physostigminum Sulphur, from which is prepared Lamellæ Physostigminæ
Dried rhizome	Extractum Pterorhizæ Liquidum, Tinctura Pterorhizæ
A neutral principle from the fruits	
Dried full grown unripe fruit	Aqua, Oleum Emulsi e
Volatile oil from unripe fruit	
Oil distilled from fresh leaves and shoots	
Dried unripe fruit	Confectio Piperis, Pulvis Opii Co
Resinous exudation from the stem	Emplastrum Piceis
Bituminous liquid obtained from the wood	Unguentum Piceis Liquid e
Dried rhizome and roots	Resina Podophylli Indici, and from it Tinctura Podophylli Indici
Dried rhizome and roots	Resina Podophylli, and from it Tinctura Podophylli
The bark	Syrupus and Tinctura Pruni Virginianæ
Dried ripe fruits	Confectio Sennæ
Heart-wood	Tinctura Lavandulæ Composita
Flower heads in powder	
Dried root	Tinctura Pyrethri
Wood of the trunk and branches	Infusum, Liquor Concentratus and Tinctura Quassæ
The inner part of the bark	Tinctura Quillaiæ and Liquor Piceis Carbonis
Salt of the alkaloid	Tinctura Quininae and Vinum Quininae
Salt of the alkaloid	Ferri et Quininae Citras, Tinct Quininae Ammon, Pulvis Quin Sulphat, Syrup Ferri Phosphoric Quin et Strych
The residue left after the distillation of the Oil of Turpentine from the crude oleo resin	Emplastrum and Unguentum Resin e Emplastrum Calefaciens, Cantharidis, Menthol, Piceis, Plumbi Iodidi, and Saponis
The erect rhizome, deprived of more or less of its cortex and dried	Extractum Rhei, Infusum Rhei, Liquor Rhei Concentratus, Pilula Rhei Co, Pulvis Rhei Co, Syrupus Rhei, Tinctura Rhei Comp
Fresh petals	Syrupus Rhinados

Bot. Name	Obtained from	Natural Order	Geographical Source
Ricini Oleum	<i>Ricinus communis</i>	Euphorbiaceæ	India
Rose Gallicæ Petala	<i>Rosa Gallica</i>	Rosaceæ	Britain and France cultivated
Rose Oleum	<i>Rosa Damascena</i>	"	Bulgaria, etc.
Rosmarini Oleum	<i>Rosmarinus officinalis</i>	Labiatæ	South of Europe, cultivated in England
Rutæ Oleum	<i>Ruta graveolens</i>	Rutaceæ	England and Southern Europe
Sabine Cacumina	<i>Juniperus Salina</i>	Conifera	Britain, Southern Europe, etc.
Saccharum Purificatum	<i>Saccharum officinarum</i>	Graminaceæ	West Indies and British Guiana
Saccharum Lactis	<i>Pos Taurus</i>	Ungulata	Domesticated everywhere
Salleinum	From various Salix, Salix fragilis and Populus	Salicaceæ	Temperate regions of the Northern Hemisphere
Sambuci Flores	<i>Sambucus nigra</i>	Caprifoliaceæ	Britain, Central and Southern Europe
Santal Oleum	<i>Santalum album</i>	Santalaceæ	India
Santonium	<i>Artemisia maritima</i> var <i>Stechmanniana</i>	Compositæ	Asiatic Russia
Sappan	<i>Casalpinia Sappan</i>	Leguminosæ	Madras and Malayan Peninsula
Sassa Radix	<i>Smilax ornata</i>	Liliaceæ	Costa Rica and South America
Sassafras Radix	<i>Sassafras officinale</i>	Lauraceæ	North America
Scammonæ Radix	<i>Convolvulus Scammonia</i>	Convolvulaceæ	Syria and Asia Minor
Scammonæ Resina	"	"	"
Scammonium	"	"	Chiefly from Smyrna, in Asia Minor
Scilla	<i>Liginea Scilla</i>	Liliaceæ	Mediterranean coasts
Scopari Cacumina	<i>Cytisus Scoparius</i>	Leguminosæ	Britain
Senegæ Radix	<i>Polygala Senega</i>	Polygalaceæ	North America
Senna Alexandrina	<i>Cassia acutifolia</i>	Leguminosæ	Soudan, imported from Alexandria
Senna Indica	<i>Cassia angustifolia</i>	"	Southern India and Arabia
Serpentaria Rhizoma	<i>Aristolochia Serpentaria</i> or <i>Aristolochia reticulata</i>	Aristolochiaceæ	Southern parts of North America
Sesami Oleum	<i>Besamum Indicum</i>	Pedaliaceæ	Indigenous to India, cultivated in Turkey and Greece
Sesum Præparatum	<i>Ovis Arlos</i>	Ungulata	Domesticated everywhere
Sinepis	<i>Brassica</i> { <i>alba</i> <i>nigra</i> }	Crucifera	Britain
Staphisagria	<i>Delphinium Staphisagria</i>	Ranunculaceæ	South of Europe
Stramonii Folia	<i>Datura Stramonium</i>	Solanaceæ	Britain, cultivated
Stramonii Semina	<i>Strophanthus "Kombé"</i>	Apocynaceæ	East Africa
Strophanthus	<i>Strychnos Nux vomica</i> and other species of <i>Strychnos</i>	Loganiaceæ	East Indies and Philippines

Parts used	Preparations into which it enters
Oil expressed from the seeds	Collodium Flexile, Lin. Sinapis, Mistura Olei Ricini, Pil. Hydrarg. Subchlor. Co.
{ Fresh and dried unexpanded petals	Confectio Rosæ Gallicæ, Syrupus Rosæ Acidum Of the Confection, Pilule Aloës Barb., Aloës et Asafetide, Aloës Socot., and Hydrarg.
Oil distilled from fresh flowers	Unguentum Aqua Rosæ
{ Oil distilled from the flowering tops	Spiritus Itosinatum, Lin. Saponis and Tinct. Lavand. Comp.
{ Volatile oil distilled from fresh herb	
Fresh and dried tops	
Juice of the sugar cane	All Syrups and Lozenges, and several other preparations.
Whey of milk, evaporated	{ Pulv. Elettarii Comp., Extract. Bellad. Alcohol., Nuclei Vom., Physostigmatis, and Strophanthi.
{ Crystalline glucoside obtained from the bark	
Flowers separated from stalks	Aqua Sambuci
Oil distilled from the wood	
A crystalline principle	Trochiscus Santonini
Heart-wood	Decoctum Sappan
Dried root	{ Extractum Sarsæ Liquidum, and Liquor Sarsæ Compositus Concentratus
Dried root	Liquor Sarsæ Compositus Concentratus
Dried root	Resina Scammonia
Resin from root	{ Pil. Scam. Co., Pulv. Scam. Co., Ext. Col. Co., Pil. Col. Co., Pil. Col. et flyoscyam
{ Gum resin obtained from living root.	
Sliced and dried bulb	{ Acetum, Oxymel, Pilula Co., Syrupus and Tinctura Scillæ; Pil. Ipecac. c. Scillâ
Fresh and dried tops	Infusum Scoparii (from dried), Succus Scoparii (from fresh).
Dried root	Infusum and Tinctura Senegæ, Liquor Senegæ Concentratus.
Dried leaflets	Confectio, Infusum, Mistura Co., Syrupus, and Tinctura Sennæ, Pulvis Glycyrrhizæ Compositus, Liquor Sennæ Concentratus
Dried leaflets	May be used in the place of Alexandrian Senna
Dried rhizome and roots	{ Infusum, Liquor Serpentinae Concentratus, and Tinctura Serpentinae, Tinct. Cnithon Co.
Oil expressed from seeds.	
Internal fat of the abdomen	Unguentum Hydrargyri
{ Dried ripe seeds of both, powdered and mixed	{ Charta Sinapis, Oleum Sinapis Volatile (from Black Mustard only), Lin. Sinapis from the Oil
Dried ripe seeds	Unguentum Staphisagriae
Dried leaves	Tinctura Stramonii
Dried ripe seeds.	Extractum Stramonii
Dried ripe seeds freed from awns	Tinctura Strophanthi and Extractum Strophanthi
The alkaloid.	

BP Name	Obtained from	Natural Order	Geographical Source
<i>Styrax praeparata</i> Succinum .	Liquidambar orientalis Pinites succinifer	Hamamelidaceae Coniferae	South-west of Asia Minor Shores of the Baltic
<i>Sumbul Radix</i> .	Fetula 'Sumbul'	Umbelliferae	Asiatic Russia
<i>Tabaci Folia</i> .	<i>Nicotiana Tabacum</i>	Solanaceae	America
<i>Tamarindus</i>	<i>Tamarindus Indica</i> .	Leguminosae	West Indies
<i>Taraxaci Radix</i> .	<i>Taraxacum officinale</i>	Compositae	Britain
<i>Terobinthina</i> <i>Canadensis</i> .	<i>Abies balsamea</i> .	Coniferae	Canada
<i>Terobinthina</i> <i>Oleum</i> .	<i>Pinus Teda, palustris</i> , and other species .	"	United States of America .
<i>Theobromatis</i> <i>Oleum</i> .	<i>Theobroma Cacao</i>	Sterculiaceae	Central America .
<i>Thuja Americannum</i>	<i>Pinus (Teda, palustris)</i>	Coniferae	{ Southern States of North America }
<i>Thymol</i> .	<i>Thymus vulgaris</i> <i>Monarda punctata</i> <i>Carum Coptum</i> .	Labiatae Umbelliferae	Manufactured in Britain .
<i>Thyroidum</i> Ste- eum	<i>Ovis Arica</i>	Ungulata .	
<i>Tinospora</i> .	<i>Tinospora cordifolia</i>	Menispermaceae	Tropical India
<i>Toddalia</i> .	<i>Toddalia aculeata</i> .	Rutaceae	Madras Peninsula
<i>Tragacantha</i>	<i>Astragalus gummifer</i>	Leguminosae	Asia Minor and Persia
<i>Turpethum</i>	<i>Ipomoea Turpethum</i>	Convolvulaceae	India and Ceylon
<i>Tylophora Folia</i>	<i>Tylophora asthmatica</i> .	Asclepiadaceae	{ Bengal, Madras Peninsula and other parts of India, and Ceylon .
<i>Ulm Cortex</i> .	<i>Ulmus campestris</i>	Urticaceae	and Southern and England .
<i>Urginea</i> .	<i>Urginea Indica</i> and <i>Scilla Indica</i>	Liliaceae .	India
<i>Uva Ursi Folia</i>	<i>Arctostaphylos Uva ursi</i>	Ericaceae	Britain
<i>Valeriana Indica</i> Rhizoma	<i>Valeriana Wallichii</i>	Valerianaceae	Himalayas
<i>Valeriana Rhizoma</i>	<i>Valeriana officinalis</i>		
<i>Vanilla</i>	<i>Vanilla planifolia</i> .	Orchidaceae	
<i>Veratri Viridis</i> Rhizoma	<i>Veratrum viride</i> .	Liliaceae	United States and Canada
<i>Veratrina</i> .	<i>Schœnocephalon officinale</i>	"	Mexico .
<i>Viburnum</i> .	<i>Viburnum pumifolium</i>	Caprifoliaceae	{ Middle and Southern United States
<i>Zingiber</i> .	<i>Zingiber officinale</i> .	Scitamineae .	{ West Indies, India, and Co- chin China .

In these Tables, Bentham and Hooker's *Genera Plantarum* has been followed in the names of the botanical sources of the drugs have been made, with the view of name is derived from the vernacular name, italics indicate that the name has been given in parentheses in the first instance where such natural orders occur.

Parts used	Preparations into which it enters.
Balsam from the trunk purified	Tinctura Benzoini Composita
Fossil resinous exudation	
{ Dried transverse slices of the root	Tinctura Sumbul
Dried leaves	
{ Fruits freed from the brittle outer part of the pericarp	Confectio Sennæ.
Fresh and dried roots	Extractum, Extractum Liquidum, Succus Taraxaci
The oleo-resin .	Collodium Flexile
Oil distilled from turpentine	Lanimentum, Lanimentum Aceticum, and Terobenum
{ Concrete oil from warm crushed seeds	All Suppositories except Glycerin
Concrete oleo resin	Emplastrum Picis
{ A crystalline substance from the oils	
{ Fresh and healthy thyroid gland	Liquor Thyroides
Dried stem	{ Infusum Tinosporæ, Liquor Tinosporæ Concentratus, Tinctura Tinosporæ
Dried root-bark	{ Infusum Toddalæ, Liquor Toddalæ Concentratus
Gummy exudation	{ Mucilago, Glycerinum, Pulv Tragac Co ; Conf Sulphuris, Pulv Opi Co, Mist Cretæ and Guaiaci, Pil Quinin, Sulph, of the Mucilage, Lotio Hydrarg Nigra
Dried root and stem	
Dried leaves	
Dried inner bark	
Younger bulbs	{ Pilulæ Ipecacuanhæ c Urgineæ, Acetum Urgineæ, Oxymel Urgineæ, Pilula Urgineæ Composita, Syrupus Urgineæ, and Tinctura Urgineæ
Dried leaves	Infusum Uvæ Ursi
Dried rhizome and rootlets	Tinctura Valerianæ Indicæ Ammoniata
Dried erect rhizome and roots	Tinctura Valerianæ Ammoniata
Dried fruit	
Dried rhizome and rootlets	
The alkaloid from cevadilla	Unguentum Veratrinæ
Dried bark	Extractum Viburni Prunifoli Liquidum
Scraped and dried rhizome	{ Syrupus and Tinctura, Zingiberis It is also used in some powders and other preparations

with respect to botanical classification, and some alterations in the method of print-making the Table more useful to students, thus quotation marks imply that the previously been used as a generic name, and an initial capital letter that the name. The names of the natural orders have been used with the termination *acae* as been retained, the name recommended by Lindley, for the sake of uniformity, has

A TABULATED COMPARISON OF THE CHIEF STANDARDISED POTENT PREPARATIONS OF THE BRITISH,
UNITED STATES, GERMAN AND FRENCH (1908) PHARMACOPŒIAS—continued

	B.P.		U.S.P.		STANDARD	B.P.	F.P.	U.S.P.	F.P.
	STANDARD	Dose	STANDARD	Dose					
Guarana									
Fluid Extract			Not less than 8% of alkali 1 grammes 35 per w/v alkali to daily multiples	30 grains = 2 grms 30 minims = 3 c.c.					
Hydrastis									
Fluid Extract			Not less than 2% of alkaline	30 grains = 2 grms					
Tincture			20 per w/v Hydr. No. 1 & 2 have 1% per w/v Hydr.	30 minims = 3 c.c. 1 fluid dram = 4 c.c.					
Hyocyanus									
Extract			0.2 per w/v Hydr. alk. in 100 w/v	4 grains = 0.25 gr.					
Fluid Extract			0.025 per w/v Hydr. alk. in 100 w/v	3 minims = 0.3 c.c.					
Tincture			0.025 per w/v Hydr. alk. in 100 w/v	30 minims = 3 c.c.					
Ipecacuanha Root									
Fluid Extract			1 per w/v Hydr. alk. in 100 w/v	1 dram = 0.065 gr.					
Tincture			1 per w/v Hydr. alk. in 100 w/v	30 minims = 3 c.c.					
Jalap									
Fluid Extract			1 per w/v Hydr. alk. in 100 w/v	1 dram = 0.065 gr.					
Tincture			1 per w/v Hydr. alk. in 100 w/v	30 minims = 3 c.c.					

A TABULATED COMPARISON OF THE CHIEF STANDARDISED POTENT PREPARATIONS OF THE BRITISH,
UNITED STATES, GERMAN AND FRENCH (1908) PHARMACOPŒIAS—continued

	B.P.		U.S.P.		P.G.W.	Dose	English 1908
	Standard	Dose.	Standard	Dose.			
Scopolia . . Extract Fluid Extract.	.	..	$\left\{ \begin{array}{l} \text{Not less than } 0.5 \\ \text{p.c. mydrate} \\ \text{alk.} \\ \text{0.05 p.c. mydrate} \\ \text{alk.} \\ \text{0.5 p.c. w/v in} \\ \text{distill. alk.} \end{array} \right\}$	$\left\{ \begin{array}{l} 0.045 \text{ gm.} = \frac{1}{2} \text{ gr.} \\ 0.01 \text{ gm.} = \frac{1}{10} \text{ grain} \\ 0.05 \text{ c.c.} = 1 \text{ minim.} \end{array} \right\}$			
			$\left\{ \begin{array}{l} \text{Not less than } 0.25 \\ \text{p.c. mydrate} \\ \text{alk.} \end{array} \right\}$	$\left\{ \begin{array}{l} 1 \text{ grain} = 0.065 \text{ gm.} \\ \frac{1}{2} \text{ grain} = 0.01 \text{ gm.} \\ 1 \text{ minim.} = 0.05 \text{ c.c.} \end{array} \right\}$			
Stramonium .							
Extract .			$\left\{ \begin{array}{l} \text{From seeds } \frac{1}{2} \text{ to } 1 \\ \text{grain} = 0.065 \text{ to } 0.065 \\ \text{gramme} \end{array} \right\}$	$\left\{ \begin{array}{l} 10 \text{ p.c. mydrate} \\ \text{alk.} \\ 0.25 \text{ p.c. w/v in} \\ \text{distill. alk.} \end{array} \right\}$			
" Fluid Extract			$\left\{ \begin{array}{l} \text{From leaves } 5 \text{ to } 15 \\ \text{minims} = 0.5 \text{ to } 0.9 \\ \text{c.c.} \end{array} \right\}$				
" Tincture							

MATERIA MEDICA

WITH

COMPOUNDS AND PREPARATIONS.

Not Official

ABSINTHIUM

WORMWOOD

The leaves and flowering tops of *Artemisia Absinthium*, L. The drug possesses an aromatic odour and a very bitter taste. It contains a crystallisable bitter principle, **Absinthin**, slightly soluble in Water, readily in Absolute Alcohol, Chloroform and Ether, also a volatile oil, to which its physiological properties are due.

Medicinal Properties—Cerebral stimulant. Absinthe, an alcoholic beverage used on the Continent, contains the chief constituents of Wormwood, its excessive use causes the disease known as absinthism.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Fr, (Absinthe), Ger (Wermut), Hung, Ital (Assenzio), Jap, Mex, Norw, Port (Losna), Rus, Span (Ajenjo), Swed and Swiss. An extract is official in Belg, Fr, Ger, Ital, Port, Russ, Span, Swed and Swiss.

Descriptive Notes—Wormwood has silvery leaves, due to the surface on both sides being covered with appressed silky hairs, each attached by a central stalk of two to four cells. The leaves are tripinnatisect near the root with short pinnules about $\frac{1}{2}$ inch long. The upper leaves are less divided, becoming trifid or simple on the inflorescence, which is a panicle of small globular greenish-yellow capitula about $\frac{1}{2}$ of an inch in diameter. It is a local plant often occurring about old farmyards, especially near the sea. The mugwort (*Artemisia vulgaris*, L.) bears a resemblance to Wormwood, but the flower heads are oblong and the leaves are dark green on the upper surface, but silvery beneath. The leaves are larger and more acute.

TINCTURA ABSINTHII—Wormwood, 1, Alcohol (60 p c), to make 10.

Dose—1 to 4 fl drin = 3.6 to 14.2 c c.

Foreign Pharmacopœias—Official in Belg, Dan, Ger, Ital, Mex, Norw, Port, Russ and Swiss, 1 in 5, Austr and Hung (compound), 1 in 10, Mex. (compound), 1 in 10, Swed and Swiss (compound), 1 in 12 $\frac{1}{2}$, all by weight.

ACACIÆ GUMMI.

GUM ACACIA

FR, GOMME ARABIQUE, GER, ARABISCHES GUMMI, ITAL, GOMMA ARABICA, SPAN, GOMA ARABICA

A colourless or yellowish product obtained from *Acacia Senegal* and other species.

Solubility.—1 in 1 of Water. Insoluble in Absolute Alcohol, Ether and Oils.

Medicinal Properties.—Demulcent. Allowed to dissolve slowly in the mouth, allays tickling cough. For a demulcent drink, 1 of Mucilage, 1 of Syrup, and 20 of Water.

Prescribing Notes.—It is chiefly used in the form of Mucilage in cough lozenges and lozenges, and to render oils, etc., emulsive with aqueous fluids.

In an 8 oz. mixture 3 dm of Mucilage of Gum Acacia are usually required for 1 oz of oils or resinous tinctures, and 10 dm for 1 oz of Balsam of Copaiba. The Mucilage should be put into a mortar and the oil added by degrees with constant trituration until an emulsion is formed, then the Water or other aqueous fluid can be added by degrees. Resinous tinctures should be added to the Mucilage which has been first diluted with twice its volume of Water, but Fixed and Volatile Oils are best added to the undiluted Mucilage. It is impossible to make a nice emulsion with Oil of Male Fern unless the Mucilage be quite fresh, in such case it is better to make the Mucilage at the time by rubbing 2 of powdered Gum with 3 of Water. Another method, which gives good results with fixed oils, is to replace the Mucilage by half its weight of powdered Gum Acacia, rub the oil with the powder, then add all at once Water equal to double the weight of the powder and rub till an emulsion is formed, now add by degrees the remainder of any aqueous liquid ordered in the prescription. Resin of Copaiba makes a nice emulsion with powdered Gum and Water. The Resin is trituated in a warm mortar, the powdered Gum mixed with it and then the Water added as in the last instance. Mucilage is used to suspend insoluble powders in mixtures, but in some cases (I mean salts, for instance) Tragacanth answers better. It used to be employed for making powders into pills, but they soon became hard, and it is now displaced by 'Dispensing Syrup' (see 'Glycerin'), Glucose, Syrup of Glucose, 'Diluted Glucose,' or Glycerin of Tragacanth.

Official Preparations.—None so Acacia; also used in the preparation of Pulvis Anagallis, Pulvis Tragacanthae Compositus, and all Trochisci.

Not Official.—Mucilage Acacie, Mistura Mucilaginis, Potio Gommeuse, Sirop de Gomme, Syrupus Acacie, also used in the preparation of Unna's Gum Pastes.

Foreign Pharmacopœias. Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Descriptive Notes.—Gum Acacia is derived from different species of *Acacia* occurring in different countries, and in some cases from more than one species in the same district, and consequently is not uniform in character. It is sorted on its arrival in European ports into different qualities for various economic uses; hence for medicinal use, only the selected or 'picked' gum corresponding to the B.P. description should be used. The finest for this purpose is the Khordofan, or 'Turkey' gum of commerce, and is derived from *Acacia Senegal*, Willd.

The Gum Acacia official in the B.P. for medicinal use is limited to the finer commercial qualities, and is characterised by the opaque outer surface, translucent interior, nearly white or faint yellowish-white colour, by readily breaking up into angular fragments, being almost odourless, with a mucilaginous but insipid taste. See also Tests.

Gum that gives a glary or ropy mucilage, like some samples of Talce and Sennaar gum, is not admissible.

Senegal gum is derived from *A. Senegal*, Delile, in the Soudan, and from *A. Adansoni*, Guill and Perr., in Senegal. It differs from the Khordofan gum in the less cracked surface and the tough and less easily fractured interior, and is characterised by the presence of

vermicular amongst the rounded pieces. The mucilage is very adhesive, and is valuable for technical purposes.

East African gum of Senegal character, and similar gum exported from Jafferabad reach England *via* Bombay, and are known in commerce as East India or Bombay gum. That exported from Senegal comes *via* Bordeaux, but all possess the same characters. True East Indian gum is very mixed in character, and several commercial varieties are recognised derived from different trees, the form and use being applied to the reddish kinds.

West African gum (1 *nilotica*, D. Lile) resembles Senegal gum, but occurs in larger pieces, without vermicular pieces intermixed.

Cape gum (1 *horrida*, Willd., and 1 *Kraussiana*, Meisn.) is distinguished by its very brittle character, and by giving a weak and not strongly adhesive mucilage.

Australian gum (1 *pycnantha*, Benth.) is usually reddish, and contains Tannin. Selected white qualities of these gums can only be used if they comply with the above tests.

The Talca or Sennar gum (1 *Fistula*, Schweinf.) that gives aropy mucilage is not easily distinguished from small gum of good quality until dissolved, except by the presence of a faint greenish tinge.

The Gummi Indicum of the *Ind* and *Col* *Ind* is derived from *Anogeissus latifolia*, Wall., and occurs in vermicular or rounded nodules, almost white if of good quality, translucent when fractured, but tough rather than brittle, and often with small fragments of bark attached, a characteristic feature which is not recognised in the Pharmacopœia. It is required, nevertheless, to yield not more than 4 p.c. of ash. The mucilage differs from that of Gum Acacia in being proportionally twice as viscid and in having a more pronounced taste.

Tests—(Gum Acacia dissolves entirely in Water, forming a distinctive more or less transparent mucilaginous liquid, which possesses a feebly acid reaction towards blue Litmus paper; it is insoluble in Alcohol (90 p.c.), its aqueous solution is precipitated by Solution of Lead Subacetate, but not by Lead Acetate Solution, and is also precipitated by strong solutions of Borax.

The more generally occurring adulterations are gums of inferior origin, Starch or Dextrin, Tannic Acid, certain Sugars, and an excessive amount of mineral matter. Inferior gums are detected by the glairy mucilage produced when the gum is dissolved in an equal weight of Water, and by the formation of a gummy deposit when this mucilage is further diluted with Water and allowed to stand. Starch or Dextrin is readily detected by Iodine Solution, Tannic Acid by the bluish-black coloration produced with Ferric Chloride Test-solution, the Sugars by Fehling's (Potassio-cupric Tartrate) Solution, and excess of mineral matter by the amount of ash left on ignition. This should not amount to more than 4 p.c. Three samples recently examined in the author's laboratory yielded 2.8, 2.9, and 3.0 p.c. The limit of ash adopted by the *B.P.* and *U.S.P.* is 4.0 p.c., by the *P.G.*, 5.0 p.c.

Preparation

MUCILAGO ACACIÆ. MUCILAGE OF GUM ACACIA

4 of washed Gum Acacia dissolved in 6 of Water, the product measures about 8½

Dose.—1 to 4 fl. dr. = 3·6 to 14·2 c.c. or more

Mucilage keeps well if made cold, poured into small bottles quite full, and stored in a cool place, but if kept carelessly it becomes sour very quickly in hot weather, and its emulsive property is impaired, if made with hot Water the change is more rapid

Incompatibles Strong Alcohol and Sulphuric Acid, Borax, Ferric salts and Lead Subacetate render it gelatinous. It is not affected by neutral Lead Acetate

Foreign Pharmacopœias.—Official in Dutch and Port., 2 and 3, Fr. and Mex., 1 and 1, Dan., Ger., Hung., Ital., Jap., Norw., Russ., Swed. and Swiss, 1 and 2, Span., 1 and 3, Belg., 1 and 9; Austr. and U.S., 31 in 100. All by weight.

Not Official.

MUCILAGO ACACIÆ (U.S.)—Washed Gum Acacia, 34; Lime Water, 31, Water, to make 100, all by weight.

MISTURA MUCILAGINOSA—Syrup, 80 minims; Mucilage of Gum Acacia, 2 fl. dr.; Water, to 1 oz. (imp.)

POTION GOMMEUSE.—Powdered Gum Arabic, 1, Simple Syrup, 3; Orange Flower Water, 1; Water, 10, all by weight.—*P.*

This has been incorporated in the *B.P.C.* as follows

Mistura Acaciæ. *Syn.* Potion Gommeuse. Gum Arabic, 1, coarse powder, 6, Syrup, 16, Orange Flower Water, 6, Distilled Water, q.s. to produce 100.

SIROP DE GOMME.—Gum, 10; Sugar, 56, Water, 31, dissolve the Gum in cold Water, then the Sugar by the aid of a water-bath, and strain.—*P.*

SYRUPUS ACACIÆ (U.S.)—Selected Gum Acacia, 10, Sugar, 80; Distilled Water, to make 100 by volume.

The formula in the previous edition of the *U.S.P.* reads.—

Syrupus Acaciæ.—Mucilage of Acacia, 25, Syrup, 75

This latter has been incorporated in the *B.P.C.*

It has been suggested that a Mucilage of Acacia should be made with Chloroform Water to overcome the tendency to fermentation.

UNNA'S GUM PASTES—A mixture of equal parts of Mucilage of Gum Acacia and Glycerol, with which are incorporated various medicaments such as Zinc Oxide and Nitric Oxide.

ACACIÆ CORTEX.—The dried bark of *Acacia arabica*, and also the dried bark of *Acacia decurrens*, Willd, the Sydney Black Wattle, or of the Victorian and Tasmanian Black Wattle, are official in *Ind.* and *Col. Add.* for India and the Australian and Eastern Colonies

Decoctum Acaciæ Corticis (1 in 16) is also official in *Ind.* and *Col. Add.* for India and the Australian and Eastern Colonies.

See also *Gummi Indicum*.

Not Official.

ACALYPHA.

The fresh and the dried herb of *Acalypha Indica*, L., are official in the *Ind.* and *Col. Add.* for India and the Eastern Colonies

Extractum Acalyphæ Liquidum (1 in 1 with Alcohol 90 p.c.), dose 5 to 80 minims = 0·8 to 1·8 c.c.

Succus Acalyphæ, the juice expressed from the bruised fresh *Acalypha* 3; Alcohol (90 p.c.), q.s. to yield 4, dose, 1 to 4 fl. dr. = 3·6 to 14·2 c.c.

Both the Liquid Extract and the Succus are official in the *Ind.* and *Col. Add.* for India and the Eastern Colonies.

ACETANILIDUM.

ACETANILIDE

C_8H_9NO , eq 134 10

B P Syn — *PHENYL ACETAMIDE* Commonly known as 'Antifebrin'

Colourless, glistening, crystalline lamellae, having a burning and somewhat bitter taste, or a white crystalline powder

It is the Monacetyl derivative of Aniline, and is prepared by the action of Glacial Acetic Acid upon Aniline

Solubility.—1 in 190 of Water, 1 in 18 of boiling Water, 1 in 12 of Alcohol (60 p c), 1 in 4 of Alcohol (90 p c), about 1 in 40 of Glycerin, it is also soluble in Ether, Benzol, and Chloroform

Medicinal Properties.—A powerful antipyretic Useful in the pyrexia of typhoid fever, erysipelas, phthisis, acute rheumatism, and smallpox An analgesic in neuralgia and other painful nerve affections, such as locomotor ataxia

In some cases it produces profuse sweating, accompanied with cyanosis and rigor, it is therefore safer to commence with small doses

From the report of a committee of the British Medical Association, it would appear that Antifebrin is less safe and less constant in its action than Antipyrine, and still less so than Phenacetin but it is possible that the ill effects noted were brought about by injudicious dosage To give it in doses of 5, 6, 8, or even 10 grains, still more to repeat these after a short interval, is highly injudicious, such doses are excessive The relative dose appears to be about one fifth that of Antipyrine (*see* Phenazonum) — *B M J* '94, i 89

Cases of poisoning by Acetanilide — death after taking 60 grains in 6 powders — *B M J N* '02, i 20, *L* '02, i 213

Antidotes.—Alcohol, Strychnine, Ether, warmth to feet, etc., Oxygen inhalation

Dose.—1 to 3 grains = 0 06 to 0 2 gramme

Ph *Gr* maximum single dose, 0 5 gramme, maximum daily dose, 1 5 grammes

Prescribing Notes.—*Best given in wafer paper or cachets, or dissolved in some weak spirit May also be suspended in Water by Compound Powder of Tragacanth or Mucilage of Gum Acacia It is sometimes given as a compressed tablet, or as an effervescent granule*

Not Official.—Mistura Acetanilidi, Pulvis Acetanilidi Compositus, Ammonol, Hydracetin, Neuronal, Phenalgin, Bromoacetanilide

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U S Not in the others

Tests.—Pure dry Acetanilide melts at 113° to 114° C (235·4° to 237·2° F), commercial Acetanilide melts at 111 36° C (232·45° F), dried Acetanilide at 112 42° C (234 35° F), and purified Acetanilide at 113 49° C. (236 28° F) The *B P* gives the melting point when dry as 113 5° C (236 5° F), the *U S P* gives 113° C (235·4° F), the *P G* 113° to 114° C (235 4° to 237 2° F.) The melting point given in the *B P*. will only be found in samples which have been purified, and dried at 100° C. (212° F) Most commercial samples melt somewhat lower It visibly softens several degrees below the actual melting point If heated below Water it fuses

considerably under 100°C . (212°F) The method of determining melting points adopted by the *B.P.* has been commented upon (*Y.B.P.*, '99, 427, *C.D.*, '99, n 219, 231) and the melting points of several substances taken by four different methods compared. Great variation was found between dried and undried commercial articles and purified products. It would appear that the *B.P.* figures are compiled and not the result of actual determination. This surmise is confirmed by a statement made by Professor Atfield that 'in the future it must be distinctly understood that the method described in the *B.P. Appendix* had not necessarily been the one by which the melting points recorded in the Pharmacopoeia had been determined.' The boiling point usually given is 295°C . (563°F), but it volatilises to a considerable extent at 100°C . (212°F), and if an aqueous solution be distilled, Acetanilide may be detected in the distillate by the Isonitrile test. The *U.S.P.* gives the boiling point as 295°C (563°F) and states that it boils without decomposition.

The distinguishing test for Acetanilide is the formation of the disagreeable and highly poisonous odour of Phenyl Isonitrile when a minute quantity of the specimen is heated with Potassium Hydroxide Solution, a few drops of Alcohol (90 p.c.) and a little Chloroform. When heated with Potassium Hydroxide Solution alone, the characteristic aromatic odour of Aniline is evolved. The test is common to the *B.P.*, *U.S.P.*, and *P.G.*, the *U.S.P.* giving quantities 0.1 gramme Acetanilide, 5 c.c. of Potassium Hydroxide Solution, and 1 c.c. of Chloroform. Various other substances yield odours somewhat resembling Phenyl-Isonitrile when treated by the Isonitrile test, but the interfering action of these substances may be destroyed by the following modification of the test, which has been tried in the author's laboratory and found to be very satisfactory, readily detecting an addition of 2 p.c. of Acetanilide. The substances experimented on were Methacetin, Phenacetin, Lactophenin, Salophen, and Phenocell Hydrochloride. The test was carried out in the following manner - A weighed quantity of 0.1 gramme of each of the substances is boiled with 10 c.c. of Water (Salophen is the only one not soluble in 10 c.c. of boiling Water), the mixture is cooled quickly by immersion in cold Water and filtered through Cotton-Wool. To 2 or 3 c.c. of the filtrate is added an equal volume of Potassium Hydroxide Solution (5 p.c.). The liquid is boiled, and small fragments of Potassium Permanganate added until the green colour first produced gives way to violet or purple. Two or three drops of a mixture made of 10 c.c. of Chloroform, 10 c.c. of Alcohol (90 p.c.), and 5 c.c. of Ammonia Solution are added, the mixture boiled, and a little more of the Chloroform, Alcohol, and Ammonia mixture added if the Permanganate has not been reduced completely. After the Chloroform has vapourised by standing a few moments, the odour is noted, and compared, if doubtful, with that yielded by a minute fragment of Acetanilide or a dilute Acetanilide solution. In testing Exalgin omit the Potassium Permanganate, otherwise the test is made as above.

The *U.S.P.* and the *P.G.* give a confirmatory test for Aniline, the

former requiring that 0.1 gramme of Acetanilide, when boiled with 2 c.c. of Hydrochloric Acid for several minutes, shall yield a clear solution which, when mixed with 3 c.c. of a 1 in 20 aqueous Phenol Solution and 5 c.c. of filtered saturated Chlorinated Lime Solution, shall produce a brownish-red colour, changing to deep blue on supersaturation with Ammonia Solution. The *BP* does not give a similar test, neither does it specifically require Acetanilide to afford 'the reactions characteristic of Acetic Acid'. The acetic radical may be satisfactorily tested for, by warming a little of the specimen with Potassium Hydroxide Solution, cooling and removing the Aniline by means of Ether. The residue left after the evaporation of the Ether from the ethereal liquid, may be examined for Aniline by the Chlorinated Lime and Ammonia test. A portion of the aqueous liquid, after the removal of the dissolved Ether, is mixed with an equal volume of Sulphuric Acid (the mixture being meanwhile kept cool), a few drops of Alcohol (90 p.c.) are added and the liquid warmed, the characteristic odour of Acetic Ether is evolved.

A cold saturated aqueous solution decolorises Bromine Water and at the same time throws down a white precipitate quite distinct even at a dilution of 1 in 2000. If the Bromine Water precipitate be dissolved by heat, it crystallises out on cooling in long tufted needles. The production of this insoluble Bromine compound distinguishes Acetanilide from Phenacetin. The Isomitate test distinguishes it from Methylacetanilide (Ezalgin), Phenacetin, and Phenazone (Antipyrine), neither of these substances yielding the reaction with this test.

The more generally occurring impurities are free Acetic Acid, unconverted Aniline, Aniline salts and mineral matter. Acetanilide should be neutral in reaction towards Litmus solution, as is also Phenacetin, but with Acetone and Aniline salts the solution becomes red, and with Phenazone blue, so that the reaction towards Litmus at once detects the presence of free Acetic Acid and affords confirmatory evidence of the presence of Acetone, Aniline salts or Phenazone. The reaction in the cold towards Ferric Chloride Test-solution ensures the absence of Acetone, Phenazone and salts of Aniline. The aqueous solution of Acetanilide should not be affected by this reagent. Acetone is a most unlikely impurity, and the object of answering a test is therefore not apparent. The value of Ferric Chloride as a reagent for Acetone is, moreover, open to question. Phenazone gives a deep red coloration, which is discharged by strong Hydrochloric Acid. On boiling, Acetanilide and Phenacetin solutions become red, and in both cases the colour is discharged by strong Hydrochloric Acid. Aniline Chloride with this reagent gives no change at first, but in a few moments becomes green. Very little importance can be attached to the behaviour of Acetanilide when boiled with Ferric Chloride Test-solution, as the Ferric Chloride itself becomes of a reddish-brown colour on boiling, owing to the formation of basic Iron salts. Acetanilide should form practically colourless solutions with Sulphuric or Nitric Acid, indicating the absence of readily charred organic impurities, Phenacetin and

Official Preparations.—Acidum Aceticum Dilutum Used in the preparation of Liquor Ammonii Acetatis, Oxyinel and Oxyinel Scillae.

Incompatibles—Ammonia, Lime, fixed Alkalis, and Carbonates.

Foreign Pharmacopœias Official in Jap, 36 p.c., sp. gr. 1.018, U.S., 36 p.c., sp. gr. 1.045 at 25° C (77° F), Norw., 30 p.c., sp. gr. 1.012, Dan., 29 p.c., sp. gr. 1.011, Dutch and Russ., 30 p.c., sp. gr. 1.011, Port. (Acido Acetico Hydratado), 48 p.c., sp. gr. 1.050, Fr., 50 p.c., sp. gr. 1.060, Swed., 25 p.c., sp. gr. 1.036.

The Acidum Aceticum of Belg., Fr., Ger., Spain and Swiss is practically Glacial, Belg. and Ger., 98 p.c., sp. gr. 1.064.

The Acidum Aceticum Dilutum of Austr., Ger., Hung. and Swiss more resembles *B.P.* Acidum Aceticum, Hung., 20 p.c., Austr., Ger. and Swiss, 30 p.c.

Tests. Acetic Acid has a specific gravity of 1.044, which corresponds with the *B.P.* figure; the *U.S.P.* gives 1.045 at 25° C (77° F), the *P.G.* Dilute Acetic Acid, which corresponds to the Acetic Acid *B.P.*, is required to possess a specific gravity of 1.041. The acid is required to contain 33 p.c. of absolute Acetic Acid, as indicated by titration with Volumetric Sodium Hydroxide Solution, 1 c.c. of an acid of the official strength neutralising 5.5 c.c. of the Normal Solution. The *U.S.P.* acid is required to contain not less than 36 p.c. by weight of absolute Acetic Acid, a weighed portion of the acid being diluted with Water and an aliquot portion of the dilution titrated with Normal Volumetric Potassium Hydroxide Solution, the *U.S.P.* stating that Phenolphthalein Test solution is to be used as an indicator of neutrality, the *P.G.* acid contains 30 p.c. w/w of pure acid. The exactly neutralised liquid yields on the addition of Ferric Chloride Test solution a deep red coloration, and on boiling the liquid, a reddish brown precipitate is thrown down; the red coloration is destroyed by Hydrochloric Acid. A portion of the acidified liquid mixed with an equal volume of Sulphuric Acid, the mixture on a while being carefully kept cool affords when warmed, after the addition of a few drops of Alcohol (90 p.c.), a characteristic ethereal odour of Acetic Ether.

The more generally occurring impurities are empyreumatic matter, Sulphurous and Formic Acids, Arsenic, Copper and Lead, Chlorides and Nitrates and fixed impurities.

All three Pharmacopœias employ the Permanganate test for fixing a limit to the quantity of empyreumatic matter; the *B.P.* and the *U.S.P.* both employ 2 c.c. of the acid previously diluted with 10 c.c. of Water. The *B.P.* directs 1 drop of Potassium Permanganate Solution (1 p.c. w/v), and requires that after the lapse of half a minute the liquid shall still retain a shade of crimson; the *U.S.P.* adds 5 drops of Tenth-normal Volumetric Potassium Permanganate Solution, and requires that the liquid shall not become entirely free from pinkish brown in less than half a minute. Both Pharmacopœias require that the Permanganate Solution shall not be immediately decolorised, the *P.G.* employs 20 c.c. of the acid and 1 c.c. of Potassium Permanganate Solution (0.1 p.c. w/w) requiring that the red coloration shall not disappear within 10 minutes. The ammoniacal Silver Nitrate test may be employed for the detection of Formic and Sulphurous Acid, 5 c.c. of the acid should yield no dark

deposit when boiled for one or two minutes with 10 c.c. of Ammonia Solution and 5 c.c. of Volumetric Silver Nitrate Solution. The *BP* uses a solution of the acid exactly neutralised with Ammonia Solution, the *PG* does not include a test. A 1 in 10 dilution of the acid slightly acidified with diluted Hydrochloric Acid Solution shall give no coloration on the addition of Hydrogen Sulphide Solution, indicating the absence of Arsenic, Copper and Lead, the *PG* includes a specific test for Arsenic requiring that 1 c.c. of the acid and 3 c.c. of Stannous Chloride Solution shall not assume a dark coloration after the lapse of an hour. The presence of Copper is also shown when the liquid is slightly supersaturated with Ammonia Solution, the liquid assuming a bluish tint if that metal be present. A 1 to 10 dilution should not yield a precipitate or turbidity with Silver Nitrate Solution or with Barium Chloride Solution, indicating the absence of Chlorides and Sulphates. It should leave no residue when evaporated to dryness, indicating the absence of fixed impurities; this requirement is common to the *BP*, and the *USP*.

Volumetric Determination.—5 c.c. require 26 c.c. of Potassium Hydroxide Solution, *Pf*, when 10 grammes are diluted with Water to 100 c.c., 50 c.c. of the dilution should require 36 c.c. of Normal *V.S.* of Potassium Hydroxide, Phenolphthalein Solution being used as an indicator, *USP*.

Preparation

ACIDUM ACETICUM DILUTUM. DILUTED ACETIC ACID

Acetic Acid, 2½, diluted with Distilled Water, *qs* to yield 20

Dose. —½ to 2 fl. drm. = 1.8 to 7.1 c.c.

Official Preparations.—Used in the preparation of Acetum Ipecacuanhæ, Acetum Scillæ, and Liquor Morphine Acetatis.

Foreign Pharmacopœias.—Official in Austr., Ger., and Swiss, 80 p.c. Acetic Acid, sp. gr. 1.041, Hung., 20 p.c., Ital., A. A. Conc. 1, Water 4, sp. gr. 1.027, Belg., A. A. 8, Water 7, Jap. and Dutch, 6 p.c., Port. (A. A. Aquoso), 10 p.c., sp. gr. 1.015, *US*, 6 p.c., sp. gr. 1.006 at 25°C (77°F), Mex., 8.68 p.c., Fr., 10 p.c., sp. gr. 1.014. See also Acetum.

Tests.—Diluted Acetic Acid has a sp. gr. of 1.006 and is officially required to contain 1.27 p.c. by weight of Hydrogen Acetate, $\text{HC}_2\text{H}_3\text{O}_2$, eq. 59.58, as ascertained by titration with Deci-normal Volumetric Sodium Hydroxide Solution. Acetic Acid *BP* is used in the preparation of the diluted acid, which is therefore naturally required to answer the tests given under Acetic Acid. The Diluted Acetic Acid of the *USP* is required to contain not less than 6 p.c. w/w of absolute Acetic Acid, Diluted Acetic Acid *PG* corresponds very closely to the Acetic Acid *BP*.

ACIDUM ACETICUM GLACIALE.

GLACIAL ACETIC ACID

Fr., ACIDE ACÉTIQUE CRISTALLISABLE, (GER., EISIGSAURE, Ital. and SPAN., see below)

A clear colourless liquid, having a pungent acetic odour. It crystallises in the cold, but again becomes fluid at temperatures above

contain 98.9 p.c w/w of absolute Acetic Acid as determined by titration with Volumetric Sodium Hydroxide Solution, 1 gramme diluted with 50 times its volume of Water neutralising 16.6 c.c. of the Volumetric Solution, the *USP* requires it to contain not less than 99 p.c w/w of absolute Acetic Acid, mentioning that Phenolphthalein Test-solution is to be used as an indicator, the *Ph* requires it to contain at least 96 p.c w/w of absolute Acetic Acid. The two latter Pharmacopœias employ Normal Volumetric Potassium Hydroxide Solution for the titration, as shown in the small type below. Both the *BP* and the *USP* require the acid to answer the tests of purity given under Acidum Aceticum, the latter Pharmacopœia includes an additional recommendation, that in carrying out the Permanganate test 2 c.c. of the acid diluted with 10 c.c. of Water should be used, that two drops of Tenth-normal Volumetric Potassium Permanganate Solution should be added and that the tint produced should not be changed to brown within two hours.

The miscibility of Turpentine with an equal volume of Glacial Acetic Acid has been proposed as a test of strength of the latter (*PJ* '99, ii. 201), but the author has shown (*PJ* '02, i. 513) that an acid conforming strictly to the *BP* titration test cannot be expected to form a clear solution with all samples of Oil of Turpentine when mixed in equal volumes. It becomes, however, a delicate test for a strength of 99.5 p.c. acid or stronger.

Volumetric Determination.—Each 5 c.c. of a mixture of 1 part Acid and 9 parts Water by weight should require at least 8 c.c. Normal Volumetric Potassium Hydroxide Solution, *Ph* 3 grammes of Acid are accurately weighed, diluted with 50 c.c. of Water, and titrated with Normal Volumetric Potassium Hydroxide Solution, Phenolphthalein T.S. being used as indicator, *USP*.

Not Official

ACIDUM ACETICUM AROMATICUM.—Glacial Acetic Acid, 72, Oil of Cloves, 9, Oil of Lavender, 6, Oil of Orange, 6, Oil of Bergamot, 3, Oil of Thyme, 3, Oil of Cinnamon, 1. All by weight, mix and filter.

This has been incorporated in the *BP C* as follows—

Acidum Aceticum Aromaticum.—Oil of Bergamot, 2.50, Oil of Cinnamon, 1.25, Oil of Cloves, 10, Oil of Lavender, 5, Oil of Orange, 5, Oil of Thyme, 2.50, Glacial Acetic Acid, *qs* to produce 100.—*BP C*

ACETUM AROMATICUM (*Gl*)—Oils of Lavender, Peppermint, Rosemary, Juniper, and Cinnamon, of each 1, Oil of Lemon, 2, Oil of Cloves, 2, Spirit, 441, Diluted Acetic Acid, 650, Water, 1900. All by weight, digest 8 days, and filter.

Preparations containing similar ingredients but in different proportions are given in Austr., Fr., Hung., Ital., Jap., Mex., Norw., Port., Russ., Swed. and Swiss.

Acetum Aromaticum.—Eau de Cologne, 980, Tincture of Benzoin, 10, Glacial Acetic Acid, 60.—*Belg*

Acetum Odoratum.—Oil of Bergamot, 0.50, Oil of Cassia, 0.10, Oil of Cloves, 0.80, Oil of Lavender, 0.20, Oil of Lemon, 0.50, Tincture of Balsam of Tolu, 1, Simple Tincture of Benzoin, 10, Alcohol (90 p.c.), 50, Glacial Acetic Acid, 4, Distilled Water, *qs* to produce 100.—*BP C*

VINAIGRE ANGLAIS.—Glacial Acetic Acid, 500, Camphor, 50, Oil of Cinnamon, 1, Oil of Cloves, 1, Oil of Lavender, $\frac{1}{2}$. All by weight, mix.

VINAIGRE DES QUATRE VOLEURS.—Tops of the Greater and Lesser Wormwood (*Artemisia Absinthium* and *A. pontica*), Rosemary, Sage,

Peppermint, Rue, and Lavender Flowers, of each 15, Calamus Root, Cinnamon Cloves, Nutmeg, and Garlic, of each 2, Camphor, 1, Glacial Acetic Acid, 15, Strong White Vinegar, 1000. Dissolve the Camphor in the Glacial Acid, macerate the other ingredients in the Vinegar for ten days, press and mix.

VAPOR ACIDI ACETICI Glacial Acetic Acid and Acetic Acid, equal parts, mix. Two teaspoonfuls in a pint of Water at 140° F. for each inhalation. Sedative and antiseptic, used for inflammatory sore throat of scarlet fever — *Throat*.

ACIDUM TRICHLORACETICUM A substitution product of Acetic Acid, but it is more antiseptic and is formed by adding Chloral Hydrate with Nitric Acid in sunlight. Colourless, deliquescent crystals, which fuse at 55° C. (131° F.), and boil at 195° C. (383° F.).

Roughly soluble in Water and in Alcohol (90 p.c.).

It is a powerful antiseptic and caustic. 1 or 2 p.c. solutions have been used as a dressing for wounds, and as a lotion and spray in acute coryza. Internally, in dilute solution, 2 to 5 grams for adults, $\frac{1}{2}$ to 1 gram for children, in gastric catarrh and summer diarrhoea — *L. M. R.* '83, 285, *T. G.* '85, 63, and '91, 319.

A test for albumen in urine — *B. M. J.* '89, ii 1114; and '90, i 681.

Foreign Pharmacopœias — Official in Dutch, Ger., Jap., Mex., Swiss, U. S.

ACIDUM ARSENIOSUM.

ARSENIOS ANHYDRIDE

B. P. Syn.—ARSENIC; WHITE ARSENIC; ARSENIOS ACID.

FR., ANHYDRIDE ARSENIQUE; GER., ARSENIIGESAURE; ITAL., ANIDRIDE ARSENIOSA; SPAN., ANHIDRIDO ARSENIOSO.

As_2O_3 , eq. 393.28.

A heavy white odourless and tasteless powder, or white opaque and crystalline or glassy and amorphous masses, obtained from arsenical ores.

Solubility.—1 in 100 of cold Water, 1 in 20 of boiling Water; 1 in 500 of Alcohol (90 p.c.), 1 in 6 of Hydrochloric Acid; 1 in 8 of Glycerin; 1 in 11 of Solution of Potash, 1 in 40 of saturated solution of Sodium Carbonate.

These figures are approximate. The published solubilities of Arsenious Acid are very contradictory, owing, no doubt, to the specimens examined being either vitreous, opaque, or a mixture of the two, and therefore of different solubilities.

Medicinal Properties.—A general tonic and alterative. Valuable in chorea, chronic (not acute) eczema, lichen, acne and psoriasis, in gout and chronic rheumatism, in painful dyspepsia, in neuralgia and spasmodic asthma, especially if anæmic or malarial in origin; in the intervals between the attacks of angina pectoris, recommended in hay fever. Given in pernicious anæmia and allied blood diseases with good result. Indispensable in all forms of weak heart accompanied by pain. Antiperiodic in malaria, in small doses it is stimulant to nervous system. In the form of **paste** it is used to destroy the pulp before stopping carious teeth.

Small doses of Arsenic, from 8 to 5 minims of Fowler's Solution, well diluted, three times a day, the best tonic treatment of the rapid heart of influenza. — *L.* '99, ii 1079.

Arsenical mixture for the removal of malignant tumours. Arsenious Acid, 1, Absolute Alcohol, 75, Aqua Dest 75, increasing the strength of Arsenic to 1 in 100, or even 1 in 80.—*B M J E '01*, n 15

Large doses in chorea. Fowler's Solution, 15 to 20 minims three times daily, reduced to 5 minims, and discontinued on the eighth day.—*B M J '01*, n 1152

Arsenic may be administered in solution, in pills, or by injection, injections are painful, Asatic pills are much used on the Continent, but on the whole the advantages are all in favour of the time honoured Fowler's Solution.—*L '03*, i 784, *B M J '01*, i 656

Dose— $\frac{1}{10}$ to $\frac{1}{4}$ of a grain = 0.001 to 0.004 gramme

Ph Ger maximum single dose, 0.005 gramme, maximum daily dose, 0.015 gramme

Prescribing Notes In solution, tablet or pill. A good pill is made by well triturating with Milk Sugar and massing with 'Diluted Glucose'. Arsenic is usually given immediately after a meal. Solution of Arsenic is frequently prescribed with Solution of Stychnium, in such cases the (acid) Liq. Arsenici Hydrochloricus should be ordered, and not the (alkaline) Liquor Arsenicalis

Incompatibles.—Salts of Iron, Magnesia, Lime Water, and vegetable astringents

Official Preparations. Liquor Arsenicalis, Liquor Arsenici Hydrochloricus. Other preparations containing Arsenium: Arseni Iodidum, Ferri Arsenas, Sodii Arsenas, Liquor Sodii Arsenatis and Liquor Arseni et Hydrargyri Iodidi

Not Official.—Liquor Ammonii Arsenitis, Pilula Asatica, Granula Dioscoridis, Arsenical Paste, Arsenical Pilula, Arsenical Cautery Powders, Leveco Water and La Bourboule Water. See also Liq. Aurii et Arseni Bromidi (*L S N F*), Sodium Cacodylicum, Arseni Bromidi Liquor

Antidotes.—The freshly prepared moist Ferric Hydroxide, or large quantities of Colicmed Magnesia, Diluted Iron followed by some Common Salt (to ensure precipitation of Ferric Hydroxide). Stomach tubes, Emetics, Mucilaginous drinks, Olive Oil, or Cotton Oil, stimulants freely, if much prostration, warmth (hot blankets and bottles)

Antidotum Arsenici (*Belg*, *Dan*, *Dutch*, *Hung*, *Ital*, *Jap*, *Port*, *Russ*, *Swiss* and *U S*)

They vary considerably in the quantities of Iron, Magnesia, and Water. *Hung*, *Jap*, *Russ*, *Swiss* and *U S* employ Ferric Sulphate; *Belg*, *Dan*, *Dutch* and *Port* use Ferric Chloride

U S formula (Ferri Hydroxidum cum Magnesi Oxido).—Mix 40 c.c. of Solution of Ferric Sulphate (sp gr 1.432) with 120 c.c. of Water, and keep the liquid in a large, well stoppered bottle. Rub 10 grammes of Magnesium Oxide with cold Water to a smooth and thin mixture, transfer this to a bottle capable of holding about 1000 c.c., and fill it with Water to about three fourths of its capacity. When the preparation is wanted for use, shake the Magnesium Oxide mixture to a homogeneous, thin magma, gradually add to it the diluted solution of Ferric Sulphate, and shake them together until a uniform smooth mixture results

Note.—The diluted Solution of Ferric Sulphate, and the mixture of Magnesium Oxide with Water, should always be kept on hand, ready for immediate use

Foreign Pharmacopoeias.—Official in *Belg*, *A* Arseniorum; *Austr*, *Dan*, *Dutch*, *Ger*, *Hung*, *Jap*, *Norw*, *Russ*, *Swed* and *Swiss*, *A* Arsenicosum; *Fr*, *Ital*, *Mex* and *Port*, Acido Arsenoso; *Spain*, *U S*, Arseni Trioxidum

Tests.—Arsenious Acid is distinguished by the following tests: (1) the microscopical appearance of the sublimate produced on heating a small quantity in a test tube, brilliant transparent octahedral crystals being formed, (2) the pale yellow coloured precipitate, soluble in Ammonia Solution and in Nitric Acid, which is thrown

down when Silver Ammonio-nitrate Solution is added to its aqueous solution, (3) when dropped upon red-hot charcoal it produces the characteristic alliacious odour of Cacodyl, and when gently heated with charcoal in a tube it is reduced, yielding a sublimate of Arsenic.

It is officially required to contain from 99.89 to 100 p.c. of pure Arsenious Anhydride as volumetrically determined by means of Deci-normal Volumetric Iodine Solution, a weighed quantity of 0.25 gramme of the acid being dissolved in Water by boiling it with five times its weight of Sodium Bicarbonate, the solution cooled, three drops of Hydrochloric Acid added to neutralise or to reconvert into Bicarbonate any Sodium Carbonate produced during the boiling. It should require from 50.8 to 50.9 c.c. of the Volumetric Solution.

The *U.S.P.* requires it to contain not less than 99.8 p.c. of pure Arsenic Trioxide, the *P.G.* not less than 98.32 p.c. of pure Arsenious Anhydride. It will be noticed from the small type below under the heading of Volumetric Determination that neither the *U.S.P.* nor the *P.G.* employ Hydrochloric Acid in performing the test. The necessity for the addition of Hydrochloric Acid is stated to be to neutralise or to reconvert into Bicarbonate any Sodium Carbonate produced during the boiling, but experiments made in the author's laboratory, using the *B.P.*, *U.S.P.*, and *P.G.* processes, showed that it made very little difference whether the Hydrochloric Acid was added or not.

The more generally occurring impurities are mineral residue, Antimony, Cadmium, Lead, Tin and Arsenious Sulphide. The specimen should be entirely volatilised on heating, indicating the absence of mineral residue. The aqueous solution acidified with Hydrochloric Acid yields with Hydrogen Sulphide Solution a lemon-yellow precipitate; it should be completely soluble in Ammonium Carbonate Solution, indicating the absence of Antimony, Cadmium, Lead and Tin. It should dissolve entirely in about ten times its weight of Ammonia Solution, forming a colourless solution when diluted with Water; it should not assume a yellow colour when acidified with Hydrochloric Acid. The *B.P.* gives no quantities, the *U.S.P.* employs 1 gramme of the substance and 10 c.c. of Ammonia Solution; the *P.G.* 1 part of Arsenious Acid in 10 parts by weight of Ammonia Solution, the test indicates the absence of Arsenious Sulphide. The *U.S.P.* has an additional test for the absence of this latter impurity, and requires that the sublimate obtained on carefully heating the substance in a dry test-tube of hard glass should not at first show a yellow colour. The *U.S.P.* states that it may be distinguished from Arsenic Acid by the lemon-yellow precipitate produced on the addition of Silver Ammonio-nitrate Test-solution to its aqueous solution, this precipitate dissolving on the addition of Ammonia Solution and depositing metallic Silver when heated.

Volumetric Determination.—A weighed quantity of 0.1 of a gramme of Arsenic Trioxide mixed with 1 gramme of Sodium Bicarbonate dissolved by the aid of a gentle heat in 20 c.c. of Water should require for decolorising not less than 20.8 c.c. of Tenth-normal Volumetric Iodine Solution, *U.S.P.*, a weighed quantity of 0.5 of a gramme of the acid dissolved in 8 grammes of Sodium Bicarbonate and 20 c.c. of boiling Water is cooled and made up to 100 c.c. 10 c.c.

of this solution should decolorise 10 c.c. of Decinormal Volumetric Iodine Solution, *P G*

Preparations

LIQUOR ARSENICALIS. ARSEFICAL SOLUTION *BP Syn* —
LIQUOR POTASSÆ ARSENIIS FOWLER'S SOLUTION

Arsenious Anhydride, 87½ grains, Potassium Carbonate, 87½ grains, Compound Tincture of Lavender, 5 fl drms, Distilled Water, q.s. to form 20 fl oz (1 in 100)

A clear red liquid possessing a Lavender odour and an alkaline reaction towards red Litmus paper. It contains 1 p.c. w/v of Arsenious Anhydride. The Liquor Potassii Arsenitis of the *USP* contains 1 p.c. w/w of Arsenic Trioxide, the Liquor Kali Arsenicosi of the *PG* contains 1 p.c. w/v of Arsenious Acid. The *Brussels Conference* recommends a standard of 1 p.c. w/w.

It is officially directed to be prepared by dissolving 87½ grains each of Arsenious Anhydride and Potassium Carbonate in 10 fl oz of Water, but it is preferably prepared by dissolving this quantity of the solid ingredients in ½ oz of the Water in a flask by the aid of heat, and diluting the solution to 10 fl oz with more of the Water, when cooled the compound Tincture of Lavender is added and the product diluted with sufficient Distilled Water to produce 20 fl oz. The *USP* employs relatively about 2 fl oz for the purpose, the *PG* also uses a minimum amount of Water to first effect solution of the ingredients, subsequently diluting with Water to the required volume. The *PG* uses simple Spirit of Lavender and the preparation is colourless.

11 minims contain 1½ grain, 1 c.c. = 0.01 gramme

Dose.—2 to 8 minims = 0.1 to 0.5 c.c.

Larger doses are given in chorea.

Ph Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes

An interesting account of the introduction of Fowler's Solution is given *CD '04*, ii 685, also in *L '01*, ii 1472.

Foreign Pharmacopœias.—Official in Aust, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Spain, Swed, Swiss and *US*, 1 Arsenious Acid in 100. All by weight.

Tests.—Liquor Arsenicalis has a specific gravity of 1.010 to 1.015. The *BP* gives no figure for the sp. gr. It is officially required to contain from 0.93 to 1.0 p.c. of Arsenious Anhydride, as determined by titration of the neutralised solution with Volumetric Iodine Solution, a faintly alkaline reaction being maintained throughout the titration by a slight excess of Sodium Bicarbonate. The *USP* dilutes 21.6 grammes of solution with Water to 100 c.c., acidifies the mixture very slightly with diluted Hydrochloric Acid, adds 2 grammes of Sodium Bicarbonate and titrates with Tenth normal Volumetric Iodine Solution, of which it is required not less than 50 c.c. should be necessary. The *PG* mixes 5 c.c. of solution with 1 gramme of Sodium Bicarbonate and 20 c.c. of Water, and after the addition of a few drops of Starch Solution adds Tenth-

normal Volumetric Iodine Solution, no permanent blue coloration should be produced by the addition of 10 c.c., but a permanent blue coloration should be produced on the further addition of 0.1 c.c. of the Volumetric Iodine Solution.

LIQUOR ARSENICI HYDROCHLORICUS. *Hydrochloric Solution of Arsenic*

Arsenious Anhydride, 87½ grams, Hydrochloric Acid, 2 fl. dm. Distilled Water, *q.s.* to form 20 fl. oz. (1 in 100)

A clear colorless liquid possessing a strongly acid reaction towards blue Litmus paper. It contains 1 p.c. w/v of Arsenious Anhydride and is thus of the same strength as *Liquor Arsenicalis*. The *U.S.P.* *Liquor Acidi Arsenosi* corresponding to this preparation contains 1 p.c. w/v of Arsenic Trioxide. The official directions for its preparation are to dissolve 87½ grams of Arsenious Anhydride and 2 fl. dm. of Hydrochloric Acid in 10 fl. oz. of Water by the aid of heat, subsequently diluting the solution to the required volume, but solution is more readily effected by dissolving the Arsenious Acid with the Hydrochloric Acid in 5 fl. oz. of Water by the aid of heat, and diluting the cool solution with sufficient Distilled Water to produce 20 fl. oz. The *U.S.P.* employs relatively about 5 fl. oz. of Water for this purpose.

11 minims contain ½ grain, 1 c.c. = 0.01 gramme, Arsenious Anhydride.

De Valangin's Solution was 4 of this strength

Dose. -2 to 8 minims = 0.1 to 0.5 c.c.

Tests.--Hydrochloric Solution of Arsenic has a specific gravity of 1.010 to 1.014, no figure for the specific gravity is given in the *B.P.* It is officially required to indicate 0.99 to 1.0 p.c. of Arsenious Anhydride as determined by the titration of 25 c.c. of the liquor with Volumetric Iodine Solution, sufficient Sodium Bicarbonate being added to ensure an alkaline reaction during titration, from 50.8 to 50.9 c.c. should be required. The *U.S.P.* employs a weighed quantity of 21.6 grammes of the solution, about 2 grammes of Sodium Bicarbonate, and 100 c.c. of Water, not less than 50 c.c. of Tenth-normal Volumetric Iodine Solution should be required.

LIQUOR ARSENII ET HYDRARGYRI IODIDI. *See ARSENII IODIDUM*

ARSENAS FERRI. *See FERRI ARSENAS.*

ARSENAS SODII. *See SODII ARSENAS.*

ARSENATIS SODII LIQUOR. *See LIQUOR SODII ARSENATIS.*

Not Official

LIQUOR AMMONII ARSENITIS is made of the same strength as *Liquor Arsenicalis*, Ammonium Carbonate being substituted for Potassium Carbonate.

PILULA ASIATICA—Arsenious Acid, ½ grain; Black Pepper, ½ grain; for one pill.

The quantities vary in different books. *Gray's Supp* gives Arsenious Acid, $\frac{1}{2}$ grain, Black Pepper, 1 grain. *Swed Fil Acid* Arsenios containing $\frac{1}{2}$ grain in each. *BPC* gives the formula which was official in *Dutch Supp* 1902, but which was omitted in *Dutch* 1905—Arsenious Anhydride, $\frac{1}{16}$ grain, Black Pepper, $\frac{1}{2}$ grain.

Used in various chronic skin diseases.

GRANULA DIOSCORIDIS (*Fr* and *Dur*)—Each granule contains 1 milligramme of Arsenious Acid.

ARSENICAL PASTE for Dentists—Arsenious Acid, 2, Morphine Sulphate, 1, Creosote, to make a stiff paste. A quantity of the size of a pin's head is ample for one application. It should be spread on Cotton Wool and placed in the tooth. It will thus destroy the sensibility of a carious tooth, and in a few hours the tooth will be ready for stopping. Cocaine has been used in place of Morphine, but it is not so good.

ARSENICAL FIBRE for Dentists—Arsenious Acid, 5, Tannin, 2, Morphine Sulphate, 5, make into a paste with Creosote, mix with Cotton Wool, and dry. This preparation is an improvement on the paste, for the latter is apt to be squeezed out over the gum edge of the cavity and cause inflammation of the surrounding tissue.

ARSENICAL PASTE (*Frère Côme's*)—For cancer, applied after the surface has been laid bare by the application of Caustic Potash. Arsenic, 1, Charcoal, 1, Red Mercury Sulphide, 4, Water, $q\ s$.

ARSENICAL CAUSTIC POWDERS Each contains from $\frac{1}{2}$ grain to $\frac{1}{4}$ grain of Arsenious Acid to 1 grain of Calomel, Vermilion or Antimony Sulphide, or of any combination of them.

La Bourboule Water contains about $\frac{1}{12}$ grain of Arsenious Anhydride in 20 fl oz.

Levico Water (strong) contains about $\frac{1}{8}$ grain of Arsenious Anhydride in 20 fl oz.

ACIDUM BENZOICUM.

BENZOIC ACID.

$\text{HC}_7\text{H}_5\text{O}_2$, eq 121.13

Fr, ACIDE BENZOIQUE, *Ger*, BENZOSÄURE, *Ital* and *Span*, ACIDO BENZOICO.

Light colourless, or almost colourless, feathery crystals, which are odourless or have a faint odour of Benzoin.

It should be preserved from the air and light in well stoppered amber tinted bottles and should be kept in a cool atmosphere.

BP and *USP* permit the use of synthetic Benzoic Acid, but *Aust.*, *Ger*, *Swiss* and *Swed* Pharmacopœias recognise only the Acid prepared from Benzoin.

The *BP* states that it is obtained from Benzoin by sublimation, or it may be prepared synthetically from Toluene, from Hippuric Acid, and from other organic compounds, from which it would appear that the authorities give a preference to the resin sublimed Acid, but their description, 'odourless when quite pure, but when obtained from Benzoin possesses an agreeable aromatic odour,' conveys an impression just the reverse of this.

The Commercial varieties of this Acid are ---

1 Resin Sublimed Acid.—Characterised by its strong empyreumatic odour, colour (varying from a pale yellow to light brown), and reducing action on both Permanganate Solution and ammoniacal Silver Nitrate, it may or may not contain Cinnamic Acid, according to the variety of the Benzoin from which it is made.

2 Resin Precipitated Acid. This is prepared from Benzoin by one of the 'wet processes,' such as boiling with Milk of Lime to form a soluble Benzoate, which is afterwards decomposed by an Acid with separation of the slightly soluble Acid Benzoin. It is practically a pure chemical, has no empyreumatic odour, and has no reducing action either on Permanganate or ammoniacal Silver solution. This is the *A. S. C. P.* only *A. S. C. P.* and is that intended to be used in the U.S., the latter, however, will pass a sufficiently purified Acid, from whatever source derived.

3 Hippuric Benzoic Acid.—When imperfectly purified this Acid retains a distinct urinous odour, and is guarded against in most Foreign Pharmacopœias, but it has been shown (*P. J.* (9) xiv, 463) that acid from this source, after resublimation, will pass the purity tests of any Pharmacopœia, so that its use is mainly a question of price.

4 Toluene Benzoic Acid.—This is manufactured in very large quantities, principally for conversion into alkali Benzoates, but partly for sale as Benzoic Acid. In the latter case it is frequently said to be sublimed over a little Gum Benzoin to give it something of the aromatic odour of the Natural Acid. This Artificial Acid conforms with most tests, but is practically certain to be contaminated with Chlorine compounds, easily detected by mixing $\frac{1}{2}$ gramme of the Acid with slaked Lime (free from Chlorine), damping with Water, igniting, dissolving the residue in Nitric Acid and adding Silver Nitrate Solution. A turbidity or precipitate is practical proof of the Toluene source of the Acid.

Solubility.—1 in 390 of Water; 1 in 12 of boiling Water, 1 in $2\frac{1}{2}$ of Alcohol (90 p.c.); 1 in $2\frac{1}{2}$ of Ether, nearly 1 in 6 of Chloroform; 1 in 12 of Benzol; about 1 in 30 of Glycerin. Borax increases its solubility in Water, 1 of Borax and 1 of Acid are soluble in 100 of Water, Sodium Phosphate also aids its solution. Soluble in aqueous solutions of the Caustic Alkalis and in hot Milk of Lime, forming Benzoates, from which it is precipitated on the addition of Hydrochloric Acid unless the solutions are very dilute.

Medicinal Properties.—Most useful in acidifying and disinfecting an alkaline and decomposing urine, a stimulant and disinfecting expectorant in chronic bronchitis and phthisis; an antipyretic in acute rheumatism.

The Sodium and Ammonium salts are preferable, as they are less irritating to the alimentary canal.

It is also useful in preventing fats from becoming rancid, and it is used as a food preservative.

Dose—5 to 15 grains = 0.3 to 1 gramme.

Prescribing Notes. *Given in cachets, in pills made up with 'Diluted Glucose' or in the form of Soda Benzoate.*

Official Preparation. *Trochiscus Acidi Benzoici*, $\frac{1}{2}$ grain in each. Contained in *Tinctura Camphore Composita*, 2 grains in each oz.; *Tinctura Opii Annoniata*, 9 grains in each oz. Used in the preparation of *Ammonii Benzoas* and *Sodii Benzoas*.

Not Official.—*Vapor Acidi Benzoici*, *Benzoic Gauze*, *Annothesm*, *Subcutin*.

Foreign Pharmacopœias. Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests.—Pure Benzoic Acid melts at 121.4°C (250.5°F), the acid obtained from Benzoin at 120°C (248°F). The U.S.P. states 121.4°C (250.5°F) and mentions that the acid sublimed from Benzoin has a lower melting point. The *P. G.* does not include a melting point. The pure acid possesses a boiling point of 249°C . (480.2°F), the acid prepared from Benzoin at 238.9°C (462°F). The carefully

neutralised aqueous solution yields on the addition of Ferric Chloride Test solution a buff coloured precipitate. The *USP* and the *PG* adopt the following method of applying this test. The *USP* adds Ferric Chloride Test-solution diluted with 2 volumes of Water to the carefully neutralised solution of Benzoic Acid in an alkali Hydroxide Solution, the *PG* mixes 0.2 of a gramme of the acid with 20 c.c. of Water and 1 c.c. of Normal Volumetric Potassium Hydroxide Solution, and after shaking vigorously filters after 15 minutes and adds 1 drop of Ferric Chloride Solution to the filtrate. In the case of this acid the *BP* does not require it to yield 'when neutralised the reactions characteristic of Benzoic Acid'. The *USP* states that when heated in a dry test-tube with 3 parts of slaked Lime, Benzene is evolved. The acid may be readily determined by titration with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, 1 c.c. of Normal Volumetric Sodium Hydroxide Solution corresponding to 0.12113 gramme of Benzoic Acid. The method for its determination in Compound Tincture of Camphor will be found under the heading of Tincture Camphorae Composita.

The more generally occurring impurities are mineral matter, Chlorobenzoic Acid, Cinnamic and Hippuric Acids. The *USP* includes a test for readily carbonisable organic impurities. Mineral matter, if present, is at once detected by a residue remaining after ignition. Chlorobenzoic Acid and Chlorine compounds (indicating when present the synthetic origin of the acid) are readily detected by the addition of Silver Nitrate Solution to the solution in Nitric Acid of the residue remaining after igniting the acid with twice its weight of Calcium Carbonate. The tests adopted by the *BP*, *USP* and *PG* are compared below in the small type paragraph headed Silver Nitrate. Potassium Permanganate and Diluted Sulphuric Acid and Potassium Permanganate Solution serve to detect Cinnamic and Hippuric Acids, and organic substances capable of reducing Potassium Permanganate Solution. The tests adopted by the *BP*, *USP*, and *PG* for the detection of these substances are described in the small type paragraph under the heading of Potassium Permanganate.

The addition of Calcium Chloride Solution to a solution of the acid rendered faintly alkaline with Ammonia Solution should produce no turbidity or precipitate, indicating the absence of Oxalic Acid. The *USP* employs Sulphuric Acid to detect readily carbonisable organic impurities.

Silver Nitrate. The solution obtained by dissolving in diluted Nitric Acid the residue left when 0.5 gramme of the Acid is heated with twice its weight of Calcium Carbonate in a closed vessel, should yield only the slightest cloudiness with Silver Nitrate Solution, indicating the absence of Chloro Benzoic Acid, *BP*. The quantities for this test given in the *PG*, and *USP* are 0.2 gramme of Acid with 0.4 gramme of Calcium Carbonate, and 0.5 gramme of Acid with 0.8 gramme of Calcium Carbonate respectively. These Pharmacopoeias direct that the Acid and Carbonate be mixed with a little Water and dried before ignition. The *PG* residue after ignition is dissolved in Nitric Acid and diluted to 10 c.c. with Water, while the *USP* residue is dissolved in 20 c.c. of Distilled Water with the aid of a slight excess of Nitric Acid, before the addition of the

test reagent. The degree of colour permitted in the *U.S.P.* is not much more than that produced by a blank experiment with the same Calcium Carbonate as used in the test.

Potassium Permanganate.—A mixture of 1 part of the Acid and 1 part of Potassium Permanganate when warmed with 10 parts by weight of Diluted Sulphuric Acid, should not evolve the odour of Benzaldehyde (Bitter Almond Oil), *R.P.* The *P.G.* and *U.S.P.* tests are made with Water instead of Diluted Sulphuric Acid, and they direct that the mixture be gently warmed for a short time (45° C (113° F) for about 10 minutes, *U.S.P.*) in a loosely stoppered test glass, and then cooled.

The colour of 2 drops of Potassium Permanganate Solution (1 p.c. w/v) should not be immediately destroyed by 0.2 grammes of the Acid suspended in 10 c.c. of Water, indicating the absence of Hippuric and Cinnamic Acids, *R.P.* A yellow to brownish cloudy solution should be obtained with 0.1 gramme Benzoic Acid and 1 c.c. Solution of Ammonia, from which Benzoic Acid is again precipitated on the addition of 2 c.c. of Diluted Sulphuric Acid; this acid mixture should almost completely decolorise 5 c.c. of Potassium Permanganate Solution (0.1 p.c.) after the lapse of 1 hour, *P.G.*

Sulphuric Acid.—On gently warming a solution of Benzoic Acid in pure cold Sulphuric Acid the colour of the solution should not become darker than light brown, and when poured into Water Benzoic Acid should separate, yielding a colourless liquid, indicating absence of readily carbonisable matter, *P.G.*

Preparation.

TROCHISCUS ACIDI BENZOICI. BENZOIC ACID LOZENGE

$\frac{1}{2}$ grain of Benzoic Acid in each, with Fruit Basis

Dose.—1 to 5 lozenges

Not Official.

VAPOR ACIDI BENZOICI.—Benzoic Acid, 3 grains, Kaolin, 12 grains; rub together and add Water, $\frac{1}{2}$ oz., Tincture of Tolu, 18 minims. Shake and make up with Water to 1 oz.—*Throat*

Extremely serviceable in sub-acute affections of the air passages.

BENZOIC GAUZE.—Contains 4 p.c. of Benzoic Acid

ANÆSTHESIN (the Ethyl-ester of ortho-ethoxybenzoic acid).—A white, odourless powder soluble 1 in 1200 of cold Water, soluble 1 in 6 of Alcohol (90 p.c.), and in ether. Introduced as a local anæsthetic and as a substitute for Orthoform (even in cases of gastric irritation).—*B.M.J.* '01, ii. 32. In suflated, dusted on or used as an ointment, is most efficient (*H.M.J.* '05, ii. 1008) in allaying the pain of burns, ulcerative stomatitis, tuberculous and malignant ulceration, whether of the larynx or other regions, or it may be given internally up to 8 grains in gastric ulcer, carcinoma, or nervous dyspepsia.

Dose.—5 to 10 grains = 0.32 to 0.65 grammes. Maximum daily dose, 40 grains = 2.6 grammes.

Suboutin (Anæsthesin paraphenol sulphonate) is a product of greater stability.

ACIDUM BORICUM.

BORIC ACID.

B.P. Syn—BORACIC ACID, HYDROGEN BORATE.

Fr., ACIDE BORIQUE; *Ger.*, BORSAURE; *Ital.* AND *Span.*, ACIDO BORICO.

H_3BO_3 , eq. 61.49.

Colourless and odourless pearly scales, or a fine, white, odourless powder, unctuous to the touch, and possessing a faintly acid and

slightly bitter taste. It volatilises in the vapour of boiling Water. It is obtained by the decomposition of Borax with a mineral acid, preferably Sulphuric Acid, or by the purification of native Boric Acid.

Solubility.—1 in 25 of cold Water, 1 in 3 of boiling Water, 1 in 4 of Glycerin, 1 in 28 of Alcohol (90 p.c.), insoluble in Ether.

Medicinal Properties.—An unstimulating local antiseptic and desiccant, it is used as a **dressing** for granulating and suppurating surfaces in general, as an **eye-wash**, 2 to 5 grains in an oz. of Water, as a **lotion, douche**, or as a **mouth-wash**, 10 to 15 grains to an oz. of Water, as a **paint** for the throat, 1 in 5 of Glycerin, as a **pessary**, 10 or 20 grains with Gelatin Mass or Oil of Theobroma.

Given internally, in cystitis associated with decomposing urine.

Used as a **dusting powder** it prevents fetid perspiration.

Small doses internally to sterilise the urine, 48 hours before operation for stricture of the urethra.—*L.* '98, i 1103.

As a **preservative**, a mixture with equal parts of Borax is more convenient than Boric Acid alone.

So called danger from the use of Boric Acid in Milk. It is far from proved that small quantities of Boric Acid, if used for a long time, are poisonous to adults or children. Large doses are, however, not considered so innocuous.—*L.* '00, i 13, 131, 574, 740.

Report of the Departmental Committee appointed to inquire into the uses of preservatives and colouring matters in foods, Boric Acid or mixtures of Boric Acid recommended to be recognised as a legal addition to cream in amount not exceeding 0.25 p.c. expressed as Boric Acid, and in butter not exceeding 0.5 p.c. expressed as Boric Acid.—*L.* '01, ii 1681, *J.S.C.I.* '01, 1228.

Report of evidence taken before the Departmental Committee. *L.* '99, ii, 1181, 1588, 1786. '00, i 207, 279, 129, 507, 586, 1409, '00, ii 276. Opinions differ as to the use of Boric Acid as a preservative of foods.—*L.* '03, i 749, 897, 920.

Influence of Boric Acid on the metabolism of children. Neither Boric Acid nor Borax in any way affected the general health and well being of the children.

B.M.J. '01, i 1397. *J.C.S.* '01, *Abstr.* ii 517.

Influence of chemical preservatives of food on health. Extremely improbable that Boric Acid if used in proper proportions would cause any injurious effect whatever to the average adult, but because of certain possible injurious effects which might be produced, the use of such preserved milk for invalids and young children is to be condemned.—*L.* '99, ii 1127, 1177.

A record of 22 cases of toxic symptoms certainly caused by Boric Acid. General conclusion is that it is capable of producing dangerous pathological effects and ought not to be considered harmless.—*L.* '01, ii 1817.

Use of Boric Acid and Borates in surgery and their internal administration, though usually free from danger, ought to be carefully guarded in patients suffering from kidney affections, and immediately discontinued should dermatitis or other toxic symptoms appear.—*L.* '99, i 23.

Skin eruptions caused by the use of Boric Acid. Attention drawn to the possibility of fomentations or ointment as sources of toxic symptoms when applied to large areas of skin. Almost all the serious cases of toxic effects of Boric Acid have occurred where it had been locally applied to an absorptive surface.—*L.* '99, i 281.

Toxic effects following the use of Boric Acid given internally, also, after irrigation with strong solutions. Erythema followed by dermatitis, which disappeared on discontinuing its use.—*B.M.J.* '99, i 17, 209, *H.M.J.* '01, ii 94, *L.* '01, ii 1511.

Dose.—5 to 15 grains = 0.32 to 1 gramme.

Prescribing Notes.—*Can be given in mixture, powders, or oachets.*

Official Preparations.—Glycerinum Acidi Borici and Unguentum Acidi Borici

Not Official.—Boric Acid dressings, Collodium Acidi Borici, Collodium Acidi Borici et Zinci Sulphatis, Lotio Acidi Borici, Mistura Acidi Borici, Pastillus Acidi Borici, Boro Glyceride, Liquor Magnesi Boratis, Magnesi Borocitras, Pulvis Acidi Borici Comp, Pulvis Magnesi Borocitricus Comp

Foreign Pharmacopœias Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., N. Am., Pers., Russ., Span., Swed., Swiss and U.S.

Tests.—Boric Acid is distinguished by its behaviour with Turmeric paper, and the green coloration which it imparts to a non-luminous flame. The Turmeric test is conveniently performed by so immersing the Turmeric paper in the fluid to be tested that only one half becomes moistened, the fluid is evaporated on a water-bath, and any brownish red colour produced is instantly noted, the paper can then be moistened with Ammonia Solution, and any further alteration in colour observed. In performing the 'flame' test, the *B.P.* uses 'Alcohol,' but it is preferable to employ purified Methylic Alcohol. The behaviour of Boric Acid towards the usual indicators of neutrality does not permit of its volumetric determination under ordinary circumstances, but in a solution containing not less than 30 p.c. of Glycerin the end reaction is quite definite with Phenolphthalein Solution. One gramme of Boric Acid should require for its neutralisation about 16.1 c.c. of Volumetric Sodium Hydroxide Solution equivalent to 99.0 p.c. of Hydrogen Borate. The *U.S.P.* has adopted the process recommended (*P.J.* '02, 1. 345, *C.D.* '02, 1. 660) for inclusion in the *B.P.*, and dissolves 1 gramme of the Boric Acid in 50 c.c. of Water after the addition of 50 c.c. of Glycerin, and states that it should require 16.2 c.c. of Normal Volumetric Sodium Hydroxide Solution, equivalent to at least 99.8 p.c. of Boric Acid. The *B.P.* does not mention a method of determination. When carefully heated Boric Acid fuses and swells up, and is finally converted into a transparent glassy hygroscopic mass. With slight modifications in the wording this statement appears in the *B.P.* and the *P.G.*, the former stating the loss of weight which should occur, namely 43.6 p.c., the *U.S.P.* states that dehydration takes place at 100° C (212° F) with formation of Metaboric Acid, and on further heating it fuses at 160° C (320° F.) to a glassy mass of Tetraboric Acid, and at a higher temperature loses all its Water and is converted into Boron Trioxide.

The more generally occurring impurities are Calcium, Copper, Lead, Iron and Magnesium, also Sulphates and Chlorides. Copper, Lead and Iron may be detected by the Hydrogen Sulphide test given in the small type below, Calcium and Magnesium by the Ammonium Oxalate or Sodium Phosphate test, Chlorides by the Silver Nitrate test and Sulphates by the Barium Chloride or Nitrate test each of which appears under its individual heading. The *B.P.* includes tests for Potassium, Sodium and Ammonium. The two former may be detected by the characteristic colours which they impart to a non-luminous flame, the latter by boiling a small quantity

of the sample with Potassium Hydroxide Solution, when no ammoniacal odour should be perceptible, nor should the issuing vapour possess an alkaline reaction towards moistened red Litmus paper. The *U S P*. includes the modified Gutzert test for Arsenic, but no special test for this substance appears in either *B P* or *P G*. A test indicating a limit of Iron is included in the *U S P* and *P G*, but not in *B P*, it is given in the small type below under the heading of Potassium Ferrocyanide. A figure of less than 10 parts per million has been suggested (*C D* '08, 1795) as a fair limit for Lead, and there seems no reason for adopting a higher limit than 5 parts per million for Arsenic.

Litmus and Turmeric.—The aqueous solution reddens blue Litmus, Turmeric paper moistened with it and dried is coloured brownish red, even in presence of Hydrochloric Acid. This brownish red colour is changed to greenish black (bluish black, *P S P*) by Alkali Hydroxide Solution (*B P* gives Potassium Hydroxide Solution, *U S P* and *P G* give Ammonia Solution).

Flame Test.—The alcoholic solution when ignited imparts to the flame a greenish tinge (especially when the solution is acidulated with Sulphuric Acid, *B P*), *B P*, *P G* and *U S P*. Glycerin solutions also colour the flame green when ignited, *P G* and *U S P*.

Barium Chloride or Nitrate.—An aqueous solution (*P G* 1:50, *U S P* 1:25) should not be affected by Barium Chloride Solution, *U S P*, or by Barium Nitrate Solution, *P G*.

Ammonium Oxalate or Sodium Phosphate.—An aqueous solution (*P G* 1 in 50, *U S P* 1 in 25) should not be affected by Ammonium Oxalate Solution, or by Sodium Phosphate Solution in presence of Solution of Ammonia, *P G* and *U S P*.

Hydrogen Sulphide.—An aqueous solution (1 in 25) should not respond to the time limit test for heavy metals, *U S P*, Hydrogen Sulphide solution should not affect an aqueous solution (1-50), *P G*.

Potassium Ferrocyanide.—The *P G* and *U S P* require that 0.5 c.c. Solution of Potassium Ferrocyanide should not immediately produce a blue colour with 30 c.c. of a 2 p.c. Solution of Boric Acid in 1 part of Hydrochloric Acid and 49 parts of Water.

Modified Gutzert Test.—5 c.c. of an aqueous solution (1 in 25) should not respond to the modified Gutzert test for Arsenic, *U S P*.

Preparations

GLYCERINUM ACIDI BORICI.

6 of Boric Acid in powder, treated with 9 of Glycerin (by weight) at 302° F. (150° C.), until the whole is reduced to 10 by weight, it is then mixed with 10 of Glycerin.

If the liquid is kept constantly stirred, instead of only 'frequently' as officially ordered, the length of time required to complete the process is considerably reduced and the product is more likely to be a good colour. Boric Acid in crystals gives rather a lighter coloured product. It is too viscid for general use.

Foreign Pharmacopœias.—Official in Russ (*Acidum Boroglycerinum*), *U S* (*Glyceritum Boroglycerini*) 5 p.c., Mex (*Glicerina Borica*) 5 p.c. Not in the others.

UNGUENTUM ACIDI BORICI. BORIC ACID OINTMENT

Finely powdered Boric Acid, 1, Paraffin Ointment, white, 9
(1 in 10)

The commercial 'Fuly subtil' contains coarse particles, before use it should be passed through a fine lawn sieve.

Foreign Pharmacopœies. Official in Austri, Dutch, Fr. (Pommade d'Acide Boric), Ger. (Neu Vaselineum Boricum, Span. and Swiss Boric Acid 1, Vaseline 9), Dan. (Boric Acid 1, Simple ointment 9), Dan. Boric Acid 1, Washed Lard 9, Dan. has also Vaselineum Boricum, Boric Acid 1, Vaseline 9, Ger. and Jap. Boric Acid 1, Paraffin ointment 9, Swed. Boric Acid 1, Wool Fat 1, Vaseline 8, U.S., Boric Acid 1, Paraffin 1, White Petroleum 8. Not in Hung., Ital., Port. or Russ.

Not Official.

COLLYRIUM ACIDI BORICI Boric Acid, 10 grains, Distilled Water, 1 oz. —*King's.*

COLLYRIUM ACIDI BORICI ET ZINCI SULPHATIS —Boric Acid, 5 grains; Zinc Sulphate, 4 grain, Distilled Water, to 1 oz. —*Middleb.*

Boric Acid, 1, Zinc Sulphate, 0.10, Distilled Water, q.s. to produce 100. —*B.P.C.*

The *B.P.C.* Supplement now employs diluted Rose Water in place of Distilled Water.

LOTIO ACIDI BORICI Boric Acid, 15 grains, Water, 1 oz. —*St. Thomas's*
Boric Acid, 9, Water, 100. —*B.P.C.*

LINTEUM ACIDI BORICI. Lint dipped in a hot saturated aqueous solution of Boric Acid and then dried. Should contain 50 p.c. of Boric Acid, and not be soaly. It is usually coloured pink.

Used as an antiseptic dressing for wounds and ulcers.

Boric Gauze, 20 p.c., Boric Wool, 25–50 p.c. —*Fr. Gaze Boriquée*, 10 p.c.

MISTURA ACIDI BORICI Boric Acid, 10 grains, Dilute Nitro-Hydrochloric Acid, 10 minims, Compound Tincture of Gentian, 1 drim., Water, to 1 oz. —*Lock.*

PASTILLUS ACIDI BORICI.—2 grains in each pastille

BORO-GLYCERIDE —A patented preparation for preserving different kinds of food. A combination of Boric Acid and Glycerin.

A solution, 1 in 20 of Water, has been used as an antiseptic in operative surgery. Used as a paint in throat affections, 1 in 2 of Glycerin, as a tampon in dysmenorrhœa.

LIQUOR MAGNESII BORATIS —*Light* Magnesium Carbonate, 1; Boric Acid, 27, Water, 127, hot and filtered. They dissolve almost completely but crystallise out when forty eight hours. Half the quantity of *Light* Colloid Magnesia can be used in the place of the Carbonate.

MAGNESII BORO-CITRAS.—A white powder, or in glistening scales, prepared by the interaction of Boric and Citric Acids with Magnesium Oxide. Stated to be useful in the uric acid diathesis and for the removal of urinary calculi —*L.* '03, 1 837, 920.

Inferior to Magnesium Lactate as a haemostatic in hæmophilia —*L.* '08, 1 96.

Dose.—15 to 30 grains = 1 to 2 grammes, several times daily.

Pulvis Magnesii Borocitratis Compositus —Magnesium Borocitrate, 1; Powdered Sugar, 2.

Dose —30 to 60 grains = 1.8 to 3.6 g.c.

PULVIS ACIDI BORICI COMP —Boric Acid, 1; Zinc Oxide, 3; Starch, 6. —*Guy's.*

Boric Acid, 24, Potassium Bromide, 24, Starch, 99, Iodoform, 2; Morphine Acetate, 1. For insufflation.

Boric Acid and Starch Powder, equal parts —*St. Thomas's* and *B.P.C.*

Listerine and **Zymocide** are liquid specialties containing Boric Acid together with other antiseptics.

Antipyrone, **Aseptin**, **Branalcan**, and **Glacialine** are preservative mixtures containing Boric Acid.

ACIDUM CARBOLICUM.

PHENOL.

C_6H_6O , eq 93.34

FR, PHENOL OFFICIAL, GER, KARBOLSAURE, ITAL, FENOLO CRISTALLIZZATO, SPAN, ACIDO FINICO

Small crystals which are colourless, but have a tendency to deliquesce and acquire a pink colour on exposure to light and air. Phenol has a characteristic odour and taste, and a strongly cauterising action upon the skin. The chief commercial source is the fraction of coal tar distilling between 150 and 200° C (302° and 392° F). It may also be synthetically prepared from Benzol, and is supplied commercially of very good quality. It should be kept in dark amber-coloured well stoppered bottles.

Carbolic Acid, or Phenol, is prepared in a crude state by treating certain oils, heavier than Water, obtained in the distillation of Coal Gas Tar, with a dilute solution of Caustic Soda, and by subsequently separating the crude Carbolic Acid from the alkaline solution by the addition thereto of a mineral Acid (usually Sulphuric). The crude Carbolic Acid thus obtained is submitted to fractional distillation and crystallisation, with other purification processes, having for their object the entire removal of the last traces of Cresylic and other Tar acids and bases, Sulphur compounds, etc.

Solubility. 1 in 13 (or a little less) of Water, 1 in 2 of Olive Oil, $3\frac{1}{2}$ in 1 of Glycerin, 3 in 1 of Chloroform, 4 in 1 of Ether, 6 in 1 of Alcohol (90 per cent), $2\frac{1}{2}$ in 1 of Benzol, $2\frac{1}{2}$ in 1 of Carbon Bisulphide, freely in Liquor Potassæ, in Liquor Sodæ, and in Volatile Oils.

The B.P. requires that at 60° F (15.5° C) 100 parts of Phenol should be liquefied by the addition of 10 parts of Water, should form a clear liquid with 30 to 40 of Water, and should be completely dissolved by 1200 of Water.

When 1 or 2 parts of melted Carbolic Acid are mixed with 1 of Water, the Acid separates on cooling in oil like globules, but when 3, 4, 5, 6, 7, 8, and even 9 of Acid to 1 of Water are mixed the solution is perfect at ordinary temperatures, when, however, the temperature sinks to 40° F or under, the 8 and the 9 will crystallise out again.

Pure Carbolic Acid readily absorbs Water from the air, and combines with it to form a definite crystalline Hydrate $2C_6H_6O.H_2O$, containing 8.74 per cent of Water and melting at 63° F (17.2° C) — *Mon.*

Medicinal Properties. — Antiseptic, disinfectant, and local anæsthetic. Given as an intestinal and gastric antiseptic in flatulence, and in dilated stomach with fermentative change, it is most efficacious in typhoid in the form of 1½ grain pills. It relieves the itching of psoriasis. It has been used with advantage internally in phthisis, bronchitis, gangrene of the lung, whooping-cough, and puerperal fever, as a prophylactic against scarlet fever. Placed in a carious tooth, or cautiously applied to the gum, relieves tooth-ache. Used as a **paint** for the throat (30 grains to 1 oz of Glycerin), as a **gargle** (2 grains to 1 oz) for tonsillitis, it used

with a **spray apparatus**, 3 grains in an oz. of Water, or for **inhalations**, 20 grains dissolved in a pint of hot Water; as an **injection** (1 grain to 1 oz. of Water), for the vagina or the bladder, as an antiseptic. Externally, used alone is a powerful caustic, as a **lotion** (15 to 30 grains to 1 oz.) for foul or syphilitic ulcers, carbuncles, scabies, ringworm and other parasitic skin diseases, (5 grains to 1 oz.) excellent for eczema and eruptions attended with itching, or as the official **ointment**. For a **mouth-wash**, see Phenate de Soude, p. 35

Carbolic Oil, 1 or 2 in 40 of Olive Oil, used for dressing scalds and burns.

Carbolic Solution, 1 or 2 in 10 of Water, used in surgery as an antiseptic

2 p.c. solutions have been used for **hypodermic injection**.

Deep hypodermic injections (4 grain to 20 minims Water) have been found most successful in erysipelas, poisoned wounds and deep seated inflammations — *Whitt*

As solutions of Carbolic Acid in strong Alcohol or concentrated Glycerin are not caustic, but become so when diluted with Water, it is suggested that in cases of burning with concentrated Carbolic Acid it would be better to remove the Acid with strong Alcohol rather than with Water. — *P. J.* '91, n. 783

Actual contact would appear to be necessary for Carbolic Acid to act as a germicide. A few inches from the surface of pure Carbolic Acid in a bottle (open to the air) putrefaction and fermentation go on as rapidly as in the open air — *P. J.* '93, n. 545.

As an ointment or plaster (1 in 15 or 20) in lupus — *M. A.* '94, 416

Carbolic Acid mixed with 5 to 10 p.c. of Glycerin injected for hydrocele — *B. M. J.* '86, i. 1164, 1214

2 p.c. spray for ... — *P. J.* '86, n. 917

Injection of a ... for anthrax — *B. M. J.* '86, n. 601; *L.* '87, n. 1186; *L. M. R.* '89, 122; *M. A.* '94, 79

1 grain in 1 oz. of Water every 4 hours for vomiting in pregnancy. — *L.* '89, i. 1121

Throat ... treated with Carbolic Acid — *B. M. J.* '97, i. 1314.

How ... pills in acute diarrhoea — *L.* '93, n. 1305

Indisputably in the treatment of tetanus, 2 minims in 30 minims of Water injected 3 times a day — *L.* '99, i. 1497, '99, n. 1589; *B. M. J. E.* '99, i. 15, '00, i. 82.

Treatment of tetanus in horses by hypodermic injections — *L.* '00, i. 538

Treatment of influenza — *L.* '99, i. 958, '00, i. 143, 509, 667, 1030

The offensiveness of the pustular stage of eruption in smallpox successfully treated by the pure liquefied acid, applied with a small camel's hair brush to the rash over a certain area of the body each day, commencing with the face and head, until the whole of the vesicles had been touched, care being taken to prevent the acid running on the healthy skin — *L.* '03, i. 518, '04, n. 1151, 1781

In the treatment of plague, 12 grains given in a mixture every two to four hours — *L.* '99, n. 1589; '00, i. 614, '01, n. 753

Fixation of movable kidney by means of strong Carbolic Acid, six cases — *L.* '03, i. 1142.

Treatment of erysipelas — *B. M. J.* '02, i. 1142.

Cases of Carbolic Acid gangrene — *T. G.* '01, 739, 192 cases of gangrene due to weak solutions of Carbolic Acid — *Med. Review* '00, 449

Treatment of various vegetations with pure Carbolic Acid — *T. G.* '01, 689.

A 5 p.c. solution failed to destroy anthrax spores after twenty four hours' exposure, but destroyed the bacillus pyocyaneus, staphylococcus pyogenes aureus, and the bacilli of typhoid fever, diphtheria, cholera, and tuberculosis — *London County Council's Report on Disinfectants*, *L.* '02, i. 754.

Carbolic Acid (5 p.c.) and Mercuric Chloride Solution (1 in 1000) proved to

be the only real germicides for tubercle bacilli—*Report of the London County Council on Disinfectants*, L '02, i 759

Poisoning from the application of Carbolic Acid to the unbroken skin A 2 p c solution being used as an application for pruritus, followed by the use on another day, after a bath, of a 4 p c solution applied to the abdomen, the thighs and the lumbar region Recovery in ten days—*L*, '02, i 1551

Alcohol stated to be one of the best antidotes in Carbolic Acid poisoning The patient is made to drink promptly a few oz of brandy, whisky, or other spirit Immediately after this a soft india rubber tube is passed through the oesophagus and into the stomach A funnel is attached to the upper end and about a pint of Water (more or less, according to the circumstances) is poured into the stomach The upper end of the tube is now depressed and the fluid is syphoned out Washing the stomach is repeated two or three times, and finally 1 drm Sodium Sulphate dissolved in a wineglassful of Water is given—*Medic's Archives* (December '99), *L*, '00, i 481, *T'G* '01, 587, '01, 707, '02, 114, *P'J* '00, i 1, '02, ii 85, *Squibb*, '01, 2563

Bacterial standardisation of disinfectants—In the 'drop' method of testing the bactericidal power of disinfectants, originated by Rideal and Walker, pure Phenol is recommended as the standard control disinfectant. It is employed in aqueous solution, and the proportion of absolute Phenol present should be determined by titration with Bromine as given under Tests The method ascertains what dilution of the disinfectant under examination kills a given bacterial culture within the same time as the standard Phenol dilution The quotient of the two dilutions indicates the efficiency of the disinfectant and is called its Carbolic Acid coefficient, or its Rideal Walker coefficient Thus if a 1 in 3000 dilution is found to be as germicidal as a 1 in 100 Phenol, the disinfectant is said to have a Phenol efficiency of 30, that is, it possesses a germicidal power thirty times stronger than Phenol In the routine testing of disinfectants the typhoid bacillus is generally used because of its medium resistance, its easy cultivation, and because it forms a good suspension, but it is noteworthy that the coefficients of a disinfectant for different bacteria are not always identical

The presence of organic matter has been found to exert an appreciable effect on the Carbolic Acid coefficients as determined by the Rideal-Walker method Meredith Blyth is of opinion that disinfectants, containing the higher Phenols, suffer great loss of efficiency when mixed with fat, albumen, faeces and urine The germicidal value of the disinfectant acting on a 'naked' organism gives little, if any, indication of its value in the presence of organic matter It appears impossible in the Phenol class to combine a low toxic value with a high germicidal value in the presence of much organic matter—*Analyst*, '06, 154.

Dose.—1 to 3 grains = 0.06 to 0.2 gramme

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme

Prescribing Notes—Usually given internally in the form of a pill 12 grains of Carbolic Acid make a good pill mass with 24 grains of *Liquorice Powder*, another good formula is, Carbolic Acid 12 grains, *Liquorice Powder* 18 grains, *Compound Tragacanth Powder* 6 grains Used in various forms as an application

Compressed tablets are supplied for extemporaneously preparing a solution

The addition of free Ammonia to solution of Carbolic Acid slowly turns the colour blue, which darkens on keeping—*P'J* (i) xxi 593

Sulphurous Acid, added in very small proportion to Carbolic Acid melted, has been stated (*C'D* '05, i 859) to counteract the tendency of the acid to acquire a red tint.

Official Preparations.—*Acidum Carboicum Liqfactum*, *Glycerinum Acid Carboici*, *Suppositorium Acid Carboici*, *Trochiscus Acid Carboici*, *Unguentum Acid Carboici* Used in the preparation of *Salol*, *Sodii Sulphocarbolas* and *Zinci Sulphocarbolas* Contained in *Injectio Ergota Hypodermica* and *Liquor Thyroidei*

Not Official—Anti catarrhal Salts, Gargarisma Acidi Carbolici, Kraus's Catheter Lubricant, Latio Acidi Carbolici, Latio Acidi Carbolici et Boracis, Liqueur Sodii Carbolatæ, Mistura Acidi Carbolici, Lind's Oil, Oleum Lubrificans, Pasta Lubrificans, Pastillus Acidi Carbolici, Resina Carbolica, Vapor Acidi Carbolici, Antiseptic dressings, Carbolic Soap, Solution of Phenate de Soude, Acidum Carboliceum Crudum, Phenolæ xl Phenol Camphor, Phenol lodatum, Pigmentum Phenol Iodati, Tribromphenol, Para mono-chlorophenol, Trichlorophenol, Sulphaminol, Sulphocarbolice Acid and Sulphocarbolates

Foreign Pharmacopœias—Aust., Belg., Dan., Dutch, Fr., Ger., Hung., Ital. (Fenolo cristallizzato), Jap., Mex. (Acido Fenico), Norw., Port., Russ., Span., Swed., Swiss and U.S.

Melting Point (Centigrade) compared with Foreign Pharmacopœias. Brit., 38.8°; Port., 35°; Dan., 39°; Fr., Spin and Swed., 42°, Hung., 36–41°; Ital., 40°; Mex., 40°; Norw., 40°; Austr., Belg., Dutch, Ger., Jap., Russ. and Swiss, 40°–42°; Swed., 39°; U.S., not lower than 40

Boiling Point (Centigrade) compared with Foreign Pharmacopœias. Austr., 178–180°; Brit., not higher than 180°; Ital., 182°; Dutch, 181–182°; Port. and Swed., b.p. not given; Fr., 180°; Belg., Ger., Jap., Norw., and Russ., 178–182°; Hung., 180–184°; Spin., 180°; Swiss, 183°–184°; U.S., not higher than 188

Antidotes.—Stomach tube, Emetics. Alcohol, Albumen, Saccharated Solution of Lime, soluble Sulphates (Magnesium or Sodium), Olive or Castor Oil; stimulants to counteract narcotism, warmth to the extremities. Hypodermic injection of Atropine Sulphate $\frac{1}{2}$ grain. Inhalations of Amyl Nitrite.

Case of Carbolic Acid poisoning by absorption treated successfully with 1 grain doses of Camphor (dissolved in Syrup) every hour for four hours—*L. M. R.* '84, 217. 100 grammes of Camphorated Oil administered in case of Carbolic Acid poisoning complete recovery—*G. D.* '99, n 1055. Recovery after swallowing 3 oz Carbolic Acid, treated by hypodermic injection of $\frac{1}{2}$ grain Apomorphine, Olive Oil and Lime Water being given freely—*L. M. J.* '89, n 1336. Soap—*L.* '89, n 415. Vinegar neutralises the effects of Carbolic Acid on the skin and mucous membrane, and is useful when Carbolic Acid has been swallowed—*L.* '96, n 251, *Pr.* Kn 220, *B. M. J.* '97, n 595.

Tests. Carbolic Acid is distinguished by the following tests: (1) its melting point, (2) the boiling point, (3) its specific gravity at the melting point, (4) the production of a deep purple-violet colour when its aqueous solution is mixed with Ferrie Chloride Test-solution, (5) the production of a white precipitate of Tribromphenol when an excess of Bromine Solution is added to its cold aqueous solution, and (6) the production of a bluish colour when 4 parts of its aqueous solution are mixed with 1 part of Anemone Solution and a few drops of Chlorinated Soda Solution, and the mixture gently warmed.

The melting point is officially required to be not lower than 102° F. (38.8° C.). The melting point 91.5° F. (33° C.) given in *B.P.* 1885, was lower than that of any other Phenol preparation, it has very properly been raised to a maximum of 102° F. (38.8° C.). It is possible with special precautions to raise the melting point of Carbolic Acid to 108° F. (42.2° C.), but the highest melting point commercially obtainable appears to be about 106° F. (41.1° C.), and no exception can be taken to a melting point of 104° F. (40° C.). The *P. G.* gives the melting point as 40° to 42° C. (104° to 107.6° F.); the *U.S.P.* requires that when Phenol is gently heated till liquid, then slowly cooled, with constant stirring until partial recrystallisation occurs, the semi liquid mass formed should have a temperature

(remaining stationary for some time) not lower than 39° C (102.2° F). A lower boiling point or a higher melting point indicates a less hydrated Phenol. The melting point and the boiling point are influenced by the presence of Water or Cresylic Acid, so that to eliminate the first it should be boiled for a few seconds and cooled. Starting with an acid melting at 104° F (40° C), 1 p.c. of added Water reduced the melting point to 98° F (36.6° C), 3 p.c. to 86° F (30° C), and 5 p.c. to 71° F (23.3° C).

The boiling point of the acid is about 180° C (356° F), the *BP* states not higher than 182° C (359.6° F), the *USP* and the *PG* 178° to 182° C (352.1 to 359.6° F). The acid has a specific gravity at its melting point of about 1.060, the *BP* states 1.060 to 1.066.

Lunge has shown (*PJ* (3) xii 593), that the addition of 1.3 p.c. of Cresylic Acid to pure Phenol reduces the melting point from 40.5° C (104.9° F) to 32.5° C (90.5° F). The lower the melting point and the higher the boiling point, the more impure is the acid. The pure acid melts at 42° C (107.6° F), and boils at 182° C (359.6° F). A useful method of judging of the purity of a commercial acid is by determining the solidifying point of the 62.5 p.c. fraction, after the first 10 p.c. fraction containing the Water and light oils has been removed.

The aqueous solution of Phenol is faintly acid to blue Litmus paper. The *BP* states that it does not immediately reddens blue Litmus, the *USP* that it is faintly acid to blue Litmus paper. An aqueous solution of Phenol yields a fine violet colour on the addition of Ferric Chloride Test-solution, the colour being pronounced even in very dilute solution. The *USP* gives quantities for this test, 1 drop of Ferric Chloride Test-solution to 10 c.c. of a 1 p.c. aqueous Phenol Solution yielding a violet-blue colour. The *PG* employs a solution of 20 parts of Phenol in 10 parts of Alcohol (90 p.c.) and states that when this solution is mixed with 1 part of Ferric Chloride Test-solution a dirty green coloration is produced, the solution when diluted with Water to 1000 parts assuming a nearly permanent light violet colour. Phenol even in dilute solution affords a white precipitate with Bromine Water. The test is common to the *BP*, *USP* and *PG*. The *USP* states that the precipitate of Tribromophenol first formed is redissolved, but becomes permanent with more of the reagent, and that when examined under the microscope it appears crystalline. The *PG* directs a 1 in 50,000 solution of the Phenol to be employed, and states that the precipitate is flocculent.

Phenol coagulates Albumen Solution and Gelatin, and forms a liquid with Camphor.

No process of assay has been introduced into the *BP*. The Eighth Decennial Revision of the *USP* has adopted the Tribromophenol or Koppeschaum's process. The Phenol is precipitated as a Bromine compound by the addition of Bromine Solution and the excess of Bromine is determined by the addition of solution of Potassium Iodide (20 p.c.), and titration of the liberated Iodine with

Tenth normal Volumetric Solution of Sodium Thiosulphate. The acid is required to show 96 p.c. of absolute Phenol. An outline of the process is given in the small type below under the heading of Volumetric Determination. The process originally recommended by Koppeschaar is given (*Zeitschrift für analytische Chemie*, xvi, 233), and consists in precipitating Phenol from its aqueous or dilute alcoholic solution with Bromine Water, the strength of the Bromine Water being determined by titration with Volumetric Sodium Thiosulphate and Potassium Iodide. The process is modified by Allen as follows. A quantity corresponding to about 0.1 of a gramme of Phenol is carefully weighed and transferred to a stoppered bottle, to this is added a solution prepared by gradually adding Bromine to 7 c.c. of Normal Volumetric Sodium Hydroxide Solution until a permanent yellow colour appears, and then boiling the liquid. When cold a measured quantity of 5 c.c. of concentrated Hydrochloric Acid is rapidly introduced, the bottle stoppered and shaken. A solution of 1.25 grammes of pure Potassium Iodide is added, the bottle shaken and allowed to stand, the liberated iodine is titrated with Deci-normal Volumetric Sodium Thiosulphate Solution, using Starch Mucilage as an indicator. In calculating out the result 7 c.c. of Normal Volumetric Sodium Hydroxide Solution neutralises 0.566 gramme of Bromine, all of which is liberated by Hydrochloric Acid. 0.1 of a gramme of Phenol requires 0.4068 gramme of Bromine, leaving a surplus of 0.1532 gramme, which would be sufficient to neutralise 19.5 c.c. of Deci-normal Volumetric Sodium Thiosulphate Solution, each c.c. of Volumetric Thiosulphate used over and above this indicates 0.00197 gramme of impurities in 0.1 gramme of the sample.

The more commonly occurring impurities are excess of Water and the presence of Cresylic Acid and Creosote. The presence of excess of Water is indicated by the lowering of the melting point, which may also indicate the presence of Cresylic Acid. The latter acid and Creosote are detected by the behaviour of a mixture of equal volumes of the liquefied acid and of the Glycerin when mixed with 3 volumes of Water. The *B.P.* and *U.S.P.* state that a clear liquid should be formed when 1 volume of Phenol liquefied by the addition of 10 p.c. of Water (8 p.c., *U.S.P.*) is mixed with 1 volume of Glycerin, and it is not rendered turbid when 3 volumes of Water are added.

Volumetric Determination.—The following process, which is a modification of that originally devised by Koppeschaar, is included in the *U.S.P.* A measured quantity of 25 c.c. (= 0.0389 gramme of Phenol) of a solution obtained by dissolving 1.566 grammes of the specimen in sufficient Water to produce 1000 c.c. is mixed in a glass stoppered bottle with 80 c.c. of Tenth-normal Volumetric Bromine Solution, 5 c.c. of Hydrochloric Acid added, followed by 5 c.c. of an aqueous Potassium Iodide Solution (20 p.c. w/v), rapidly introduced; the mixture is shaken, the stopper and neck of the bottle rinsed with a little Water, allowing the washings to run into the bottle, 1 c.c. of Chloroform added and the mixture well shaken. The liberated iodine is titrated with Tenth-normal Volumetric Sodium Thiosulphate Solution, of which the number of c.c. used subtracted from 80 and the remainder multiplied by 4 yields the percentage of absolute Phenol present in the specimen operated upon.

Preparations

ACIDUM CARBOLICUM LIQUEFACTUM. LIQUEFIED PHENOL
LIQUEFIED CARBOLIC ACID

Phenol, 10, Distilled Water, 1, by weight

It forms a clear, colourless, highly refractive liquid possessing the characteristic odour of Phenol. It has a tendency to acquire a pinkish tint, and should therefore be preserved in well-stoppered dark amber tinted glass bottles

Dose.—1 to 3 minims = 0.06 to 0.2 c c

Foreign Pharmacopœias—Official in Austr, Belg, Fr, Ger, Hung, Jap and Russ, Carbolic Acid, 100, Water, 10. Dan, Ital., Norw, Span, Swed and Swiss, Carbolic Acid, 90, Water, 10. Dutch, Carbolic Acid, 100, Water, 20, U S, not less than 86.4 p c. Not in the others

A weak **Solution of Carbolic Acid** is official in the following Pharmacopœias—Ital (Aqua Phenicata), and Port (Aqua Phenica), 1 in 100, also 1 in 1000, (Aqua Carbolisata), Austr, Belg, Dutch and Ger, 1 in 50, Hung (Aqua Carbolata), Mex (Soluciones de Acido Fenico), and Swiss (Aqua Phenolata), 1 in 100, Dan (Solutio Phenoli), Fr (Soluté de Phénol), Norw (Solutio Acidi Carbolici) and Swed (Solutio Phenoli), 1 in 50, Span (Aqua Fenicada), 1 in 50. The *Brussels Conference* adopted a strength of 2 p c w/w for Phenol: Solutio seu Aqua Phenolata

Tests—Liquefied Carbolic Acid has a specific gravity ranging from 1.064 to 1.069, a boiling point not higher than 182° C (359.6° F), and it should answer the tests of identity and purity given under Acidum Carbolicum

When a small quantity of solution (say, 2 fl dm, in a test-tube with a thermometer dipping into the solution) is cooled to about 10° C (50° F) and gently stirred, it becomes a mass of crystals, which will entirely disappear when the temperature rises to 14.4° C (58° F)

GLYCERINUM ACIDI CARBOLICI. GLYCERIN OF PHENOL

Phenol, 1, Glycerin, sufficient to produce 5 (1 in 5)

Used as an antiseptic packing in cases of acute middle ear catarrh, and has given good results—*B M J* '04, n 1210. A 10 p c aqueous solution of the acid with Cocaine affords relief of pain in non suppurative middle-ear disease—*B M J* '04, n 1211

Mixed with an equal bulk of Water, may be applied to aphthous stomatitis, or to ulcers in the month, or to inflamed tonsils. Mixed with 20 or 30 parts of Water, it makes an excellent gargle

Foreign Pharmacopœias—Official in U S, 1 in 5, Mex, 1 in 50; Port, 1 in 100, Span, 1 in 100. Not in the others

SUPPOSITORIA ACIDI CARBOLICI. PHENOL SUPPOSITORIES

Each suppository contains 1 grain of Phenol, 2 grains of White Beeswax, and about 12 grains of Oil of Theobroma

The addition of Wax to Oil of Theobroma raises the melting point of the mass without producing the required firmness. Suggested that future *B P* basis should be Theobroma alone, *I. J.*, '00, n 85

TROCHISCUS ACIDI CARBOLICI. PHENOL LOZENGE

1 grain of Phenol in each, flavoured with Tolu.

Dose.—1 to 3 lozenges

UNGUENTUM ACIDI CARBOLICI. PHENOL OINTMENT OINTMENT OF CARBOLIC ACID—*B.P.* '85.

Phenol, 1, Glycerin (by weight), 3; white Paraffin Ointment, 21 (1 in 25)

In *B.P.* '85 ointment, part of the Phenol crystallised on keeping, and acted as a caustic. To avoid this the Phenol is now dissolved in Glycerin.

Attention has been directed (*L.* '05, i. 514) to the tendency of the *B.P.* Ointment to crystallise, and a modified formula suggested.—Phenol, 1, Hard Paraffin, 6; Soft Paraffin, 18. This formula is stated to be superior to the official one, and to be recommended especially for obstetric use. The desirability of adopting alternative formulas for ointment bases in the *B.P.* Ointment bases may be classed into (1) Non absorbent ointment bases, e.g., Hard and Soft Paraffin, (2) Emollient absorbent ointment bases, e.g., Lard and Olive Oil, (3) Systematic absorptive dermatine bases, e.g., Wool Fat.

The ointment made according to the directions contained in the British Pharmacopoeia has been the subject of a good deal of discussion from time to time, some authorities holding that even now it is not a satisfactory product, whilst others hold that it is. According to *P.J.* '05, i. 344, the formula as it stands has been well designed, and it produces an excellent ointment.

The *B.P.* 1885 ordered Carbolic Acid 1, Soft Paraffin 12, Hard Paraffin 6 = 1 in 19. The *Companion* noted the fact that the Carbolic Acid did not dissolve until the melted mixture was heated to 60°C (140°F) or over, and that part of the Carbolic Acid crystallised on keeping the ointment, but no crystals were formed when the strength was reduced to 1 in 30. The *U.S.P.* has now been altered to 8 in 100, and this has been incorporated in the *B.P.C.* under the title **Paraffinum Carbolisatum**.

Foreign Pharmacopœias Ointment Fr (Pommade de Phénol), Phenol 1, Vaseline 99, Ital (Pomata Fenata) Carbolic Acid 1, Benzoylated Lard 99; Mex (Pomada de Acido fenico), Carbolic Acid 1, Alcohol 1, Vaseline 98; U.S. (Unguentum Phenolis), Phenol 3, White Petroleum 97. Not in the others.

Not Official.

ANTI-CATARRHAL SALTS—Phenol, 1, Eucalyptus Oil, 1, Pumilio Pine Oil, $\frac{1}{2}$, Strong Iodine Solution, $\frac{1}{2}$; Camphor, 1, Ammoniated Alcohol, 2; Pine Sawdust, 2, or *q.s.*—*Martindale*

This is incorporated in the *B.P.C.* under the title **Vapor Eucalypti Compositus** with *q.s.* anti-catarrhal Salts.

GARGARISMA ACIDI CARBOLICI.—Glycerin of Phenol, 1 fl. oz.; Water, to 1 pint.—*St. Thomas's*

This has been incorporated in the *B.P.C.*

Glycerin of Carbolic Acid, 25 minims; Water, to 1 fl. oz. *London.*

LOTIO ACIDI CARBOLICI.—Carbolic Acid, 30 grains; Water, 8 oz. This lotion applied to mosquito bites relieves the itching, pain, and swelling. If mixed with a little Glycerin and sponged over the face and hands before retiring to rest, the mosquitoes will not bite until the Acid be thoroughly evaporated by the heat of the skin—*L.* '78, ii. 280.

See also Foreign Pharmacopœias under 'Acidum Carboliceum Liquefactum'

Lotio Acidi Carbolici.—Carbolic Acid (crystals), 1 oz., Water, to 20 fl. oz.—*London*

Liquefied Carbolic Acid, 21 minims; Water, to 1 oz.—*Westminster*

Phenol Crystals, 1, Water, to 20. Usually diluted with Water before use, as an antiseptic in surgery—*B.P.C.* See also Carbolic Solution, p. 28.

LOTIO ACIDI CARBOLICI ET BORACIS—Glycerin of Phenol, 2 fl oz, Glycerin of Borax, 2 fl oz, Water, to 1 pint To be diluted with five to ten parts of Water—*St Thomas's*

This has been incorporated in the *B P C*

Lotio Boracis cum Acido Carbolico.—Bicarbonate of Sodium, 20 grams, Borax, 20 grams, Glycerin of Carbolic Acid, 1 fl dm, Distilled Water, to 1 oz—*Royal Free*

Lotio Acidi Carbolici cum Borace—Glycerin of Carbolic Acid, 6 minims, Glycerin of Borax, 6 minims, Water, to 1 oz—*Children's Hospital, G O S*

MISTURA ACIDI CARBOLICI (Roth)—Pure Carbolic Acid, 12 minims, Tincture of Iodine, 16 minims, Tincture of Orange, 90 minims, Syrup, 3 dm, Water, to 8 oz Recommended for use in typhoid fever, 1 oz every 4 hours—*J. '88, 1 1214*

LUND'S OIL—Phenol, 1, Castor Oil, 4, Almond Oil, 20—*Joel*

Phenol, 1, Castor Oil, 4, Almond Oil, 15—*Companion* (1899).

This has been incorporated in the *B P C* under the title **Oleum Lubrificans**.

A solution of Carbolic Acid in Oil is frequently used to lubricate and at the same time disinfect catheters, but Koch's experiments show that such a solution has no antiseptic power, and they ought to be first disinfected with an aqueous solution, and afterwards oiled—*Brunton*

Kraus's Catheter Lubricant—Tragacanth, 2 5, Glycerin, 10, Phenol Water (3 p c), 90

This paste facilitates the passage of the catheter and is easily washed off in warm Water—*P J '99, n 529*

This has been incorporated in the *B P C* as under—

Pasta Lubrificans Syn Catheter Paste—Carbolic Acid, 3, Glycerin, 10, Tragacanth, 2 50, Distilled Water, q s to produce 100—*B P C*

PASTILLUS ACIDI CARBOLICI—Carbolic Acid, $\frac{1}{2}$ gram, Glycerogelatin, 18 grains in each

RESINA CARBOLICA—Resin, 4, Carbolic Acid Crystals, 4, Chloroform, 3—*R D H*

Resin, 45, Carbolic Acid, 85, Chloroform, 20—*B P C*

VAPOR ACIDI CARBOLICI—Pure Carbolic Acid, 420 grams, Water, 1 dm, dissolve 20 drops in a pint of Water at 140° F for each inhalation Antiseptic, very serviceable in syphilitic and carcinomatous ulcerations

CARBOLIC ANTISEPTIC DRESSINGS—Absorbent Wool and Lint containing 5 and 10 p c of absolute Phenol, Gauze, 5 p c Tow, 5 p c, Ligatures; Protective Oiled Skin, Silk Sutures Fr (Gaze Phénolée), 2 to 5 p c, Belg, Gauze, 5 p c, Jap, Cotton Wool, 5 p c, Ital, Gauze, 5 p c, Wool, 2 p c, Mex, Gauze, 10 p c, Stupa Carbolisata (*B P C*), Carbolic Acid, in crystals, 5, Methylated Ether (0 720), 100, Jute Tow, dried, 95, Stupa Carbolisata Composita (*B P C*), Jute Tow, dried, 85, Tar, 1, Carbolic Acid, 10, Methylated Ether, 100

CARBOLIC SOAPS—These contain 10 p c and 20 p c of Phenol

SOLUTION DE PHENATE DE SOUDE—Phenol, 100, Solution of Caustic Soda (sp gr 1 332), 20, by weight, Water to measure, 1000 *P*

One part of this solution to 80 of Water makes a good antiseptic mouth-wash.

Liquor Sodii Carbolatis—Phenol, 8, Caustic Soda, 3 $\frac{1}{2}$, Distilled Water, 100—*Martindale*

Liquor Sodii Carbolatis. Syn. Solution of Sodium Phenate—Carbolic Acid, 8, Sodium Hydroxide, 4, Glycerin of Cochineal, 1, Distilled Water, q s, to produce 100.—*B P C*

A formula is given (*A J P '90, 109*) as representing the proprietary article sold under the name 'Phénol Sodique' Coal tar, 2 troy oz, Soda, 120 grams, Water sufficient to make 20 fl oz

ACIDUM CARBOLICUM CRUDUM A yellowish, yellowish-brown, or reddish-brown liquid having a strongly empyreumatic and disagreeable odour. It consists chiefly of **Cresylic Acid** (see p. 12) and is largely used for disinfecting drains, etc.

Foreign Pharmacopœias.—Official in Hung., Ital., Jap. and Russ. Not in the others.

PHENOSALYL.—A specialty containing Phenol and Salicylic Acid introduced as an antiseptic.

PHENOL-CAMPHOR—Carbolic Acid and Camphor will form a liquid in any proportion between Camphor 3, Carbolic Acid 1 and Camphor 1, Carbolic Acid 3, but most a still better apper to use in excess of Camphor. The formula $C_{10}H_{16}O$, attributed to this compound, corresponds with molecular weights of each, Carbolic Acid and Camphor (Carbolic Acid 2 parts, and Camphor 3 parts).

A colourless or faintly liquid with an odour of Camphor. Soluble in Alcohol (90 p.c.), Ether, Chloroform, and Oils. Insoluble in Glycerin and in Water.

Used as a local anæsthetic for toothache—*T. G.* '85, 269; *L.* '89, n. 8077. Camphor, 60, Phenol, 19, Water, 1, is not so caustic as Carbolic Acid—*Pr.* xl. 128, and xlv. 52.

Acidum Carbolicum Camphoratum (Hager)—Camphor, 3, Carbolic Acid, 1.

This has been incorporated in the *B. P. C.*

Camphora Carbolisata (Hager)—Camphor, 25, Carbolic Acid, 9; Spirit, 1.

Carbolic Acid 1, Camphor 3, has been applied in diphtheria, etc., either pure or mixed with an equal volume of Oil of Almonds.

PHENOL IODATUM (Iodised Phenol **Pigmentum Phenol Iodati**)—Iodine, 40 grains, Liquefied Carbolic Acid, 1 cc.—*Hoep. Women, and Samaritan.*

Applied on a dressed wound or forceps in chronic endometritis and endocervicitis, with or without a previous curetting. A fl. dr. diluted with 20 oz. of Water is used as a vaginal douche in midwifery—*L.* '88, n. 862.

Iodine, 10, Liquefied Carbolic Acid, to make 100—*B. P. C.*

Pigmentum Iodi Carbolicum.—Iodine, 1; Liquefied Phenol, 4.—*Guy's.*

Pigmentum Iodi Carbolisatum.—Iodine, Potassium Iodide, and Phenol, of each 4 grains (Glycerin, $\frac{1}{2}$ oz., Water, to 1 oz.—*Central Throat*

This is sometimes used at half strength.

It has been incorporated in *B. P. C.* as follows—Iodine, 1, Potassium Iodide, 1; Phenol, 1; Glycerin, 50; Distilled Water, q. s. to produce 100.

TRIBROMPHENOL (Bromol)—White crystalline powder, with a slightly aromatic odour. A sample melted at $185^{\circ} F$ ($85^{\circ} C$).

Solubility.—1 in 2 of Alcohol (90 p.c.), 1 in 1 of Ether; 1 in 2 of Chloroform, almost insoluble in Glycerin, &c. Dissolves in Caustic Alkaline Solutions, 1 in 200 of Glycerin, 1 in $7\frac{1}{2}$ of Olive Oil.

It possesses considerable antiseptic properties.

PARA-MONOCHLOROPHENOL—Occurs in crystalline needles. Soluble in Alcohol, Ether, and Fixed Oils, but practically insoluble in Water. It possesses a stronger microbicidal power than Phenol, but its employment requires careful watching—*H. M. J. E.* '95, i. 11; *J. E.* '95, n. 551; '98, i. 61; *C. D.* '95, i. 224.

5 or 10 p.c. Glycerin solution in laryngeal phthisis, by intralaryngeal injection, also $\frac{1}{2}$ to $\frac{1}{4}$ p.c. solutions for inhalations. Under the name of **Menthosol**, a mixture of Menthol and Para-chlorophenol in 5, 10 and 15 p.c. solutions has been introduced—*B. M. J. E.* '02, i. 43.

TRICHLORPHENOL.—White crystalline powder, with a pungent, somewhat tarry odour.

Solubility.—1 in 1 of Alcohol (90 p.c.); 2 in 1 of Ether; 1 in $1\frac{1}{2}$ of Chloroform, 1 in 1000 of Water, 1 in 9 of Glycerin; 1 in 8 of Olive Oil.

It forms salts with Ammonium, Potassium, Magnesium, Calcium and Lead.

It is stated to be an antiseptic and deodorant much stronger than Carbolic Acid.

SULPHAMINOL (Thio oxydiphenylamine) Yellow, odourless, tasteless, powder Insoluble in Water, soluble in Alcohol and Ether Antiseptic dusting powder Internally in doses of 3 to 4 grains = 0.2 to 0.36 gramme, three or four times a day in cystitis

SULPHOCARBOLIC ACID ($\text{H C}_6\text{H}_4\text{SO}_3$) — Phenol para sulphonic Acid is formed by the action of Sulphuric Acid upon Carbolic Acid when warm Phenol-ortho sulphonic Acid is produced in the cold

A few years ago it was revived under the name **ASEPTOL**, a syrupy liquid, mixing in all proportions with Water, Alcohol, and Glycerin

AMMONIUM, MAGNESIUM, POTASSIUM, and SODIUM SULPHOCARBOLATES all crystallise in tufts of acicular crystals more or less white, **COPPER SULPHOCARBOLATE**, in transparent light blue interlacing prisms, the **IRON** salt, in small brown micaceous crystals, the **ZINC** salt, in tabular crystals

The Sodium and Zinc Sulphocarbulates are official See **SODIUM SULPHOCARBOLATES** and **ZINC SULPHOCARBOLATES**

ACIDUM CHROMICUM.

CHROMIC ANHYDRIDE

CrO_3 , eq 99.38

FR, ACIDE CHROMIQUE CRISTALLISE, GER, CHROMSÄURE, LIQF, ANHYDRIDE CHROMICA, SPAN, ACIDO CROMICO

Small purplish red crystals, which are slightly hygroscopic even when absolutely free from Sulphuric Acid, but much more so when a trace of the latter is present They possess a strong corrosive action on animal and vegetable tissues

It should be kept in well stoppered, dark amber-tinted glass bottles

It is produced by the action of Sulphuric Acid upon Potassium Bichromate

Solubility.—About 2 in 1 of Water, Alcohol decomposes it

It is a powerful oxidising agent, and is liable to cause sudden combustion or *explosion* in contact with strong Alcohol, Ether, Glycerin, and some other organic matters

Medicinal Properties.—Disinfectant, antiseptic, deodorant It is a powerful caustic (1 in 1 of Water), and is used by means of a pointed glass rod, great care being taken to protect the adjacent parts by plaster or ointment, having moist lint ready to absorb any superfluous Acid, 100 grains to 1 oz Water is used to remove warts, lupus, and condylomata, 1 in 40 of Water may be applied to ulcers of mouth or pharynx, and 1 in 2000, or even 4000, is used as a lotion for putrid sores, leucorrhœa and ozæna

It is of great importance for its use as a caustic that Chromic Acid should be free from Sulphuric Acid

A warm concentrated solution rapidly dissolves all animal tissues

5 p.c. Solution of Chromic Acid applied with a brush to the feet after bathing gave excellent results in the German Army as a remedy for excessive perspiration — *P J* (8) xx 504

The pure Acid fused on the point of a probe has been applied with success to nasal mucous membrane in cases of hay fever and paroxysmal sneezing — *M A* '94, 817

Official Preparation Liquor Acidi Chromici

Not Official Garg. acidi Chromici, Latio Acidi Chromici and Pigmentum Acidi Chromici

Foreign Pharmacopœias Official in Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex. (Acid. Chromico), Norw., Port., Russ., Span., Swed., Swiss and U.S. (Chromii Trioxidum).

Tests.—Chromic Acid is distinguished by its melting point, which should be from 192° to 193° C. (377·6 to 379·4 F), the *B.P.* gives 192° C. (377·6 F), the *U.S.P.* 192° to 193° C. (377·6 to 379·4 F), whilst the *P.G.* does not include a melting point, the production of green coloured liquids when its aqueous solutions are mixed with reducing agents, the evolution of Oxygen when strongly heated, and the evolution of Chlorine when warmed with Hydrochloric Acid. It liberates Iodine from Potassium Iodide Solution, and this reaction has been utilised by the *U.S.P.* as a means of determining the percentage of Chromic Anhydride. The Iodine liberated when the specimen is treated with an excess of Potassium Iodide, in a solution acidified with Hydrochloric Acid, being titrated with Tenth-normal Volumetric Sodium Thiosulphate Solution; it is required to contain not less than 90 p.c. of pure Chromium Trioxide.

The more generally occurring impurity is Sulphuric Acid, which is tested for, in an acidified solution, by Barium Chloride Solution, only a slight opalescence should be afforded.

Barium Chloride or Nitrate.—An aqueous solution 1 in 50 (1 in 100 *P.G.* and *U.S.P.*) previously acidulated with Hydrobromic Acid. Should be unaffected by Barium Chloride Solution. (*P.G.* uses Barium Nitrate Solution) This test is common to *B.P.*, *P.G.* and *U.S.P.*, the *P.* test permits a slight opalescence.

Volumetric Determination—A weighed quantity of 1 gramme is dissolved in 100 c.c. of Water. A measured quantity of 8·8 c.c. of this solution is mixed with 2 c.c. of Hydrochloric Acid and about 1 gramme of Potassium Iodide and diluted with 100 c.c. of Water. The Iodine liberated should require not less than 22·5 c.c. of Tenth-normal Volumetric Sodium Thiosulphate Solution to decolorise it, using 5 c.c. of Starch Test Solution as an indicator. 1 c.c. of Tenth normal Volumetric Sodium Thiosulphate indicates 4 p.c. of pure Chromium Trioxide, *U.S.P.*

Preparation.**LIQUOR ACIDI CHROMICI.**—SOLUTION OF CHROMIC ACID.

Chromic Anhydride, 1; Distilled Water, 3.

It forms an orange-red caustic liquid, possessing an acid reaction. It is officially required to contain the equivalent of 25 p.c. of Chromic Anhydride, CrO_3 , or 29·5 p.c. of Chromic Acid, H_2CrO_4 .

Foreign Pharmacopœias Official in Belg. and Fr., Chromic Acid, 1, Distilled Water, 1, dissolve.

Tests.—The specific gravity is officially stated to be 1·185; but a solution prepared by dissolving 10 grammes of Chromic Anhydride (free from Sulphuric Acid) in 30 c.c. of Distilled Water had a sp. gr. of 1·214. As the official Chromic Acid is used in its preparation, it is naturally required to answer the tests of identity and purity given in the monograph on this Acid.

Not Official

GARGARISMA ACIDI CHROMICI—Chromic Acid, 1 grain, Water, to 1 oz.—*Lock*

This has been incorporated in the *B.P.C.* as follows—Chromic Acid, 1, Distilled Water, *q s* to produce 500

LOTIO ACIDI CHROMICI—Chromic Acid, 10 grains, Water, 1 oz.—*University and Westminster*

PIGMENTUM ACIDI CHROMICI—Chromic Acid, 10 grains, Water, to 1 oz. In chronic superficial glossitis and secondary syphilis.—*Throat*

ACIDUM CHRYSOPHANICUM.

See CHRYSAROBINUM

Not Official**ACIDUM CINNAMICUM**

$C_9H_8O_2$, eq 146.95

There are two varieties of Cinnamic Acid (1) Medicinal, (2) Artificial

Medicinal Cinnamic Acid is obtained from natural Cinnamic Acid derivatives, *e.g.*, *Styrax*, etc. Colourless, glistening crystals, having a faint, fragrant odour. Sparingly soluble in Water, soluble in Alcohol (90 p.c.), and in Ether. It is converted into Benzaldehyde on oxidation with Potassium Permanganate. Used in the form of intravenous or intermuscular injection in pulmonary tuberculosis. In 5 p.c. alcoholic solution as an application in laryngeal tuberculosis.

Artificial Cinnamic Acid is prepared synthetically by the interaction of Benzaldehyde and Acetylchloride. Colourless crystals, sometimes possessing a faint odour of Benzaldehyde. Sparingly soluble in Water, readily soluble in Ether and Alcohol. Its use is limited to the preservation of solutions, dressings, etc.

See also Sodii Cinnamas, p 1112

ACIDUM CITRICUM.

CITRIC ACID

$H_3C_6H_5O_7$, H_2O , eq 208.50

FR, ACIDE CITRIQUE, GER, CITRONENSAURE, ITAL AND SPAN, ACIDO CITRICO

Large colourless crystals, or a white crystalline powder, possessing an acid taste. Obtained principally from Lemon Juice, which may contain from 5 to 8 p.c.

Solubility—10 in 6 of Water, and measures 12½; 1 in 2 of Glycerin, 10 in 15 of Alcohol (90 p.c.), 1 in 8 of Ether, almost insoluble in Benzol and Chloroform.

The solubility of Citric Acid in Ether naturally varies with the amount of Alcohol and Water which the Ether contains. The above figure represents its solubility in Ether (sp. gr. 0.735). The figure for Ether Purus (sp. gr. 0.720) is 1 in 40.

Medicinal Properties.—Refrigerant and sialagogue, relieves thirst in fevers. Efficacious in scurvy, for which it is also prophylactic.

Citric Acid 1, dissolved in Distilled Water 12½ (or 35 grains in 1 oz.) is a substitute for Lemon Juice, but does not keep long without spoiling

17 grains of Citric Acid neutralise about	{	24½ grains Potassium Bicarbonate.
		20 " Potassium Carbonate
		20½ " Sodium Bicarbonate.
		31½ " Sodium Carbonate.
		12½ " Ammonium Carbonate
		11½ " Magnesium Carbonate.

Dose.—5 to 20 grains = 0.32 to 1.3 grammes

Prescribing Notes.—Usually given in powders to be taken with each dose of an alkaline mixture during effervescence; or in solution, directing the quantity to be taken with the alkaline mixture

Incompatibles.—Potassium Tartrate, alkaline Carbonates, Acetates, and Sulphides.

Official Preparations.—Used in the preparation of *Liquor Ammonii Citratis*, *Liquor Potassii Citratis*, *Liquor Sodii Citratis*, *Infusum Citri*, *Infusum Citri et Ammonii Citratis*, *Infusum Citri et Potassii Citratis*, *Infusum Citri et Sodii Citratis*, *Infusum Citri et Ammonii Citratis Effervescentum*, and in all the granular effervescent Citrates.

Not Official.—*Syrupus Acidii Citrici*.

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests.—Citric Acid is distinguished by the following tests (1) the production of a white precipitate insoluble in solution of Potassium Hydroxide, but soluble in Ammonium Chloride Solution and in solution of alkali Citrates, when its neutralised solution is boiled with Calcium Chloride Solution; (2) the white precipitate soluble in Ammonia Solution, produced when the neutralised solution is treated with Silver Nitrate Solution. In contradistinction to Tartrates no mirror is produced when this ammoniacal solution is warmed. The melting point of Citric Acid is rather a variable figure (*P.J.* [3], xxi 1051) The fully Hydrated Acid melts at about 70° C. (158° F.), and the anhydrous acid at 153° C. (307.4° F.), but the crystals, and more particularly the powder, begin to dehydrate even below 70° C. (158° F.), so that intermediate figures will be obtained according to the manner in which it is heated. No melting point is given in the *B.P.*; the *U.S.P.* gives between 152° and 153° C. (305.6° and 307.4° F.), and states that at about 75° C. (107° F.) it begins to lose Water of crystallisation and becomes anhydrous at about 135° C. (275° F.).

It is officially required to indicate 99.38 p.c. of Hydrogen Citrate as ascertained by titration with Volumetric Solution of Sodium Hydroxide, Phenolphthalein Solution should be employed as an indicator of neutrality, Litmus Solution not being suitable; the *U.S.P.* requires it to contain not less than 99.5 p.c. of pure Citric Acid, the *P.G.* does not state a requisite percentage.

The more generally occurring impurities are Lead, Tartaric Acid, and mineral matter. Calcium, Iron, heavy metals, Oxalic and Sulphuric Acids may also be present. The most important and most likely impurity is Lead, and the official method of testing for

this metal has given rise to severe criticism. The *BP* directs the solution to be nearly neutralised before the addition of Hydrogen Sulphide and omits reference to limiting the quantity of Hydrogen Sulphide Solution to be employed. *PG* directs the solution to be nearly neutralised, whilst the *USP* makes the solution acid by the addition of a few drops of Hydrochloric Acid, and then requires that it shall not respond to the time limit test for heavy metals. The methods employed by the Pharmacopœias for testing for Tartaric Acid are also different, the *BP* adopting the tests with Ferrous Sulphate and Hydrogen Peroxide, and with Ammonium Molybdate and Hydrogen Peroxide, requiring that an aqueous solution of the acid should not afford a purple or violet coloration when supersaturated with Potassium Hydroxide Solution after the previous addition of a little Ferrous Sulphate Solution and a few drops of Hydrogen Peroxide Solution. The Ammonium Molybdate test is performed by mixing 1 gramme of the acid with 5 c.c. of Ammonium Molybdate Solution, and adding a few drops of Hydrogen Peroxide Solution. The test is not entirely satisfactory, as other substances besides Tartaric Acid yield a similar coloration, notably metallic particles, such as Lead, the *USP* adopts the test with Solution of Potassium Acetate. The mineral residue left on incineration with free access of air should according to the *BP* and *USP* not amount to more than 0.05 p.c. and according to *PG* 0.5 grammes should leave no weighable residue.

A standard of 5 parts per million for Lead and 1 part per million for Arsenic has been suggested (*CD* '08, 1795).

Potassium Acetate Solution—1 gramme of the powdered Acid dissolved in 5 c.c. of Solution of Potassium Acetate (1-3) should remain clear even after the addition of an equal volume of Alcohol (absence of Tartaric or Oxalic Acid). *USP*

Pure Sulphuric Acid—1 gramme Citric Acid with 10 c.c. pure Sulphuric Acid should become at most yellow in colour but not brown, when warmed in a test tube on a water-bath for 1 hour, *PG*

Ammonium Oxalate Solution—A 10 p.c. w/w aqueous solution should not be affected by Ammonium Oxalate Solution, *PG*, 5 c.c. of a 1 in 10 aqueous solution nearly neutralised with Ammonia Solution should remain clear on the addition of 1 c.c. of Ammonium Oxalate Test Solution, *USP*

Barium Chloride or Barium Nitrate—A 10 p.c. w/w aqueous solution should not be affected by Barium Nitrate Solution, indicating the absence of Sulphates, *PG*. 10 c.c. of a 1 p.c. w/w aqueous solution after the addition of a few drops of Hydrochloric Acid should not be rendered turbid within 5 minutes on the addition of 1 c.c. of Barium Chloride Test Solution, indicating limit of Sulphuric Acid, *USP*

Volumetric Determination—34.75 c.c. of a solution of 5 grammes Citric Acid in Water to measure 100 c.c., should require not less than 24.87 c.c. of Normal Volumetric Potassium Hydroxide Solution using Phenolphthalein Test Solution as indicator, which is equivalent to not less than 99.5 p.c. of pure Citric Acid, *USP*

Not Official.

SYRUPUS ACIDI CITRICI. *Syn.* Syrupus Citri:

Belg.—Citric Acid, 20, Syrup, 950, Water, 20, Spirit of Lemon, 2, Alcohol (94 p.c.), 8

Fr.—Citric Acid, 10, Syrup, 970, Alcoolature de Citron, 20

Hung.—Citric Acid, 2, Sugar, 100, Water, 50

Mex.—Citric Acid, 10, Simple Syrup, 370, Water, 20

Port.—Citric Acid, 1, Syrup of Lemons, 98, Water, 1

Russ.—Citric Acid, 3, Syrup, 150, Librosacchari Citri, 1.

Swed.—Citric Acid, 1, Syrup, 19

Swiss.—Citric Acid, 2, Spirit of Lemon, 1 5, Water, 2 5, Simple Syrup, 94

U.S.—Citric Acid, 10, Water, 10; Tincture of Fresh Lemon Peel, 10; Syrup, to make 1000

B.P.C.—Citric Acid, 3; Tincture of Lemon, 3; Syrup, *q. s.* to produce 100

All by weight except U.S. and *B.P.C.*

Not Official.

ACIDUM CRESYLICUM.

CRESYLIC ACID. CRESOL.

C_7H_8O , eq. 107.25.

A colourless or slightly yellow liquid, with a tarry odour, obtained from Coal tar. It should be preserved in well stoppered glass bottles of a dark amber colour.

There are three isomeric Cresols, but the principal constituent of the 'crude Carbohc Acid' of commerce (the source of commercial Cresylic Acid) is the 'Para cresylic Acid, with more or less of its isomers.—*Allen*.

A mixture of the three was introduced as an antiseptic under the name of **Trikresol**.

By the same process which yields Salicylic Acid from Phenol, the three isomeric Cresols yield three corresponding Cresotic or Cresotinic Acids, the Sodium salts of which have been used in Medicine.

Solubility.—1 in 80 of Water, and mixes in all proportions with Alcohol (90 p c.), Ether, Chloroform, Glycerin, and Olive Oil.

Medicinal Properties—Disinfectant and antiseptic. Used as an **inhalation** in whooping-cough, and other respiratory affections.

Prescribing Notes.—*It is far less soluble in Water than Carbohc Acid, and therefore not so convenient.*

Foreign Pharmacopœias—Official in Fr, Ger, Jap, and Swiss, *Cresolum crudum*, a yellowish liquid, Austr, *Kresolum*, acicular crystals which become yellow or brown on keeping, soluble in .38 parts of Water. Austr has also *Kresolum Liquifactum* (kresol, 100, Water, 10), and *Aqua Kresolica* (about 1 of liquefied kresol in 50). Belg., *Cresolum crudum*, yellow or brownish liquid, Dutch, *Cresolum crudum*, yellow, yellowish brown, or reddish-brown liquid, Mex, *Cresilol*, colourless fluid, Span, *Cresol*, yellowish-red liquid, U.S., *Cresol*, colourless or straw coloured liquid.

Tests.—Cresol has a specific gravity of 1.045 to 1.048, and a boiling point when pure of 203° C. (397.4° F.), but a good commercial sample may boil 10° C. lower. The *U.S.P.* specifies a sp gr. at 25° C. (77° F) of 1.036 to 1.038 and a boiling point from 195° to 205° C (383° to 401° F). Neither sp gr nor boiling point are given in *P.G.* It does not crystallise at the freezing point of Water. Its aqueous solution gives a transient blue colour with solution of Ferric Chloride

The converse of the test for Cresol (Cresylic Acid) in Phenol applies here. Equal volumes of Cresol and Glycerin should form a clear solution from which on the addition of three volumes of Water most of the Cresol should separate

The more generally occurring impurities are hydrocarbon oils.

These are readily detected by mixing equal volumes of Cresol and Sodium Hydroxide Solution (10 p c) which should form an almost clear liquid from which on standing no appreciable oily layer shall separate

Sodium Hydroxide Solution—1 c c of Cresol should dissolve in 1 c c of a 10 p c solution of Sodium Hydroxide leaving no appreciable liquid residue, *U S P*, when a volume of 10 c c of Cresol is shaken in a 200 c c stoppered graduated measure, with 50 c c of a 15 p c aqueous solution of Sodium Hydroxide and 50 c c of Water only a few flocks shall separate out. On the subsequent addition of 80 c c of Hydrochloric Acid and 10 grammes of Sodium Chloride and shaking, an oily layer collects on the top of the liquid, when allowed to remain at rest, which shall amount to 8.5 to 9 c c, *P G*

A similar test to the above is official in the *P Belg*

Ferric Chloride Solution—When 0.5 c c of Cresol is shaken with 300 c c of Water a liquid is obtained which is coloured bluish-violet by the addition of solution of Ferric Chloride—*Belg* and *P G*

LIQUOR CRESOLIS COMPOSITUS—Cresol, 50, Linseed Oil, 35, Potassium Hydroxide, 8, Water, *qs* to make 100, all by weight—*U S P*

Germicidal value stated to be greater than Carbolic Acid—*L '07*, ii 544, '08, i 576

Cresylic Acid, by weight, 50, Linseed Oil, by weight, 35, Potassium Hydroxide, 8, Alcohol, 4, Distilled Water, to produce by weight 100 *B P C*

LIQUOR CRESOLI SAPONATUS (*Ger*, *Jap* and *Swed*)—(Crude Cresol, 1, Sapo Kalinus (*vi Sapo*), 1 warm and mix to form a yellowish-brown fluid

A similar mixture is official in *Belg* and *Swiss*, under the title of Cresolum saponatum

The liquor Cresoli Saponatus of the Dutch Pharmacopœia is the same as above, but the product is finally made up to 2 with Water, and **Lysol** is given as a synonym

Aqua Cresolica (*Belg* and *Ger*)—Cresol Soap Solution, 1, Distilled Water, 9.

Cresol Soap Solution, 6, Water, 94—*Jap*

Liquor Cresolis Glycerinatus—Cresol, 50, Linseed Oil, 13, Potassium Hydroxide, 4.25, Alcohol, 2, Glycerin, 6, Distilled Water, *qs* to produce 100, all by weight—(*W J Uglow Woolcock*) *P J '07*, ii 334

This has been incorporated in the *B P C* under the title **Solutio Cresolis Saponatus**, with synonym as above

The following is understood to be the composition of the various proprietary preparations—

JEYES' FLUID—A preparation of Tar Oil containing 20 p c. The cresol saponified with resin and alkali. It forms a permanent emulsion with Water

Used in 1 or 2 p c solution, and for the same purposes as Carbolic Solutions. An injection of 1 in 400 is excellent in gonorrhœa and œzema, and in obstetric practice on account of its hæmostatic as well as its antiseptic properties. It is useful as an ointment in erysipelas

PEARSON'S ANTISEPTIC is a similar preparation to Jeyes' Fluid. The Carbolic Acid coefficient for *B. Typhosus* is 2.6—*Public Health, Dec 1903*

ARTMANN'S CREOLIN—A solution of Tar hydrocarbons in Sulphocresylic Acid. It forms a turbid liquid with Water

EUROPHEN (Di isobutyl orthocresol Iodide)—A fine light, brownish-yellow, amorphous powder, having an aromatic saffron like odour. Introduced as a substitute for Iodoform. Insoluble in Water or Glycerin, freely soluble in Absolute Alcohol, Chloroform or Ether. Applied as a dusting Powder, or 10 p c Ointment.

Losophan (Tri iodometacresol), a white or yellowish white powder, insoluble in Water, soluble 1 in 7 of Alcohol (10 p c), 1 in 4 of Ether, 1 in 6 of

Chloroform, and **Traumatol** (Iodo-resol), are compounds of Cresol and Iodine, introduced into medicine chiefly as substitutes for Iodoform.

LYSOL.—Sp. gr. 1.047. A transparent brown syrupy liquid, which forms a clear solution with Water. It is a solution in neutral Soap, of Tar Oils which distil between 187° and 200° C., and are present to the extent of about 47 p.c.

Foreign Pharmacopœias.—Official in Dutch, Mex. and Russ.

Injection of 9 to 12 c.c. of a 1 p.c. solution of Lysol into the spinal canal in cerebro-spinal meningitis. — *L.* '02, ii. 1188.

SAPROL.—Tar Oils dissolved in large excess of Hydrocarbons. Inflammable.

SOLUTOL. Sodium Cresolate in excess of Cresol, powerfully disinfectant, but caustic, and not intended for surgical purposes.

SOLVEOL.—Cresols in Sodium Cresolate, soluble in Water. Non caustic, and used for surgical purposes.

METAKALIN. A readily soluble solid preparation of Cresol. Supplied in two forms (a) in cartridges of 24 drms each, (b) in the form of tablets of 15 grains each, each tube containing ten tablets.

CRESOTINIC ACID.—There are three varieties of Cresotinic Acid, the ortho-, meta-, and para-. The only one of these which has received attention as a disinfectant is the para-cresotinic acid.

PARA-CRESOTINIC ACID occurs in long white needles or in rhombic prisms. Antiseptic, antirheumatic and antipyretic.

SODIUM PARACRESOTINATE.—The Sodium salt of Para-cresotinic acid. A white crystalline powder. Employed in doses and for purposes similar to Sodium Salicylate.

A Calcium Cresotinate is also known as a disinfectant.

Not Official.

ACIDUM FORMICUM

Formic Acid, Amebic Acid

H_2CO_2 , eq. 4 to 67.

A clear, colourless, volatile liquid, possessing an irritating odour and strongly acid taste. It is miscible in all proportions with Water.

It is a powerful stimulant of muscular action, retards fatigue, and gives a markedly increased capacity for work. In its tonic effects it is closely allied to Kola, Coca and Caffeine. It has also a diuretic effect, but not to the same extent as Theobromine. In small doses it markedly improves the appetite and general nutrition.

Dose.—2 to 5 minims = 0.12 to 0.3 c.c. in aerated Water.

The acid has been given (*B.M.J.* '08, ii. 56) in doses of from 8 to 10 drops, taken four times daily in a little Vichy or aerated Water.

It has been used in the treatment of lupus, and in cancer in the form of a subcutaneous injection in doses of 0.1 to 1.0 c.c. of a solution 1 to 100,000 to 1 in 1000, according to the age and condition of the patient.

Three hundred cases of diphtheria treated with 5 to 20 minims of 25 p.c. solution of Formic Acid 4-hourly for 10 to 14 days, plus antitoxin, the death rate from cardiac failure reduced from 86 to 2 p.c.—*Edin. Med. Jour.* '06, ii. 887.

No drug so good in chorea, as it steadies the muscular system without weakening it.—*L.* '07, ii. 1886.

Formates are stated (*L.* '05, i. 892; '07, i. 1170) to increase the power of resistance to fatigue and to promote energy. They are also slightly diuretic. 3 grammes (45 grains) of the Potassium or Sodium salt may be given daily, or a gramme and a half (23 grains) of the Lithium salt. The treatment is generally continued for ten days, followed by an interval of ten days' rest.

The acid has been given (*B M J E* '05, ii 39) in rheumatic conditions in the form of a 2½ to 3 p c solution, eight drops as an injection after injecting five to eight drops of a 1 p c solution of Cocaine as a local anæsthetic, and has been administered (*J*, '05, ii 907) in doses of 4 grammes of the normal solution in the treatment of tremor. The remedy may also prove useful in certain forms of chorea.

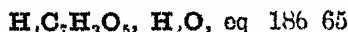
Foreign Pharmacopœias.—Official in Ger and Swiss

Tests.—Formic Acid has a specific gravity of 1.060 to 1.068. A white crystalline precipitate is produced when it is mixed with Lead Acetate. When either the acid itself or its neutralised solution is warmed with Silver Nitrate Solution a precipitate of metallic Silver is thrown down, when warmed with Mercuric Oxide metallic Mercury is precipitated, with Mercuric Chloride a white precipitate of Calomel is produced. On the addition of Ferric Chloride Test solution a reddish-brown solution is produced, which on heating throws down a reddish brown precipitate, when warmed with concentrated Sulphuric Acid, Carbon Monoxide gas is evolved which burns with a blue flame. The percentage of acid may be determined by titration with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, 5 c c of Formic Acid should require 28 to 29 c c of the Volumetric Solution, indicating 24 to 25 p c w/w of absolute acid. When diluted with Water and acidified with Nitric Acid, it should yield no immediate precipitate or turbidity with Silver Nitrate Solution, indicating the absence of Chlorides, when neutralised with Ammonia Solution it should neither yield a precipitate nor a turbidity on the addition of Calcium Chloride Solution, nor a coloration on the addition of Hydrogen Sulphide Solution, indicating the absence of Oxalic Acid and of Lead and Copper. When heated it should be entirely volatilised without leaving a weighable residue.

ACIDUM GALLICUM.

GALLIC ACID

TRIHYDROXYBENZOIC ACID



FR, ACIDE GALLIQUE, GER, GALLUSSAURE, ITAL, ACIDO GALLICO, SPAN, ACIDO AGALLICO

White or light brownish-yellow crystalline needles or prisms, odourless, and possessing an astringent and faintly acidulous taste.

It is produced by the hydrolysis of Tannic Acid, Sulphuric Acid being the acid generally employed for this purpose. It is also present in small proportion in Galls. It should be preserved in dark amber-tinted glass bottles.

Solubility.—1 in 100 of cold Water, 1 in 3 of boiling Water, 1 in 8 of Alcohol (90 p c), 1 in 50 of Ether, 1 in 6 of Glycerin with heat. Gallic Acid 1, and Potassium Citrate 1, will dissolve in 30 of Water.

Medicinal Properties. It was at one time used as a local astringent, but it is far inferior to Tannic Acid for this purpose. As Tannic Acid is converted into Gallic Acid in passing through the circulation, the latter has been given for the purpose of arresting hæmorrhage in remote vessels, but is now generally believed to be useless in such cases.

Useless for pulmonary or renal hæmorrhage.—*B M J* '00, ii 1070

Dose.—5 to 15 grains = 0.32 to 1 gramme.

Prescribing Notes.—With twice its weight of Sugar, may be taken three times a day in Water, in powders or in cachets. It is also given in pills 30 grains of Acid and 30 grains of Gum Tragacanth will make 6 pills.

Incompatibles. Spiritus, Etheris Nitrosi, metallic salts

Not Official.—Gallanol, Galloformin, Gallobromol.

Foreign Pharmacopœias.—Official in Belg., Fr., Ital., Jap., Mex., Port., Span., Swi., and U.S. Not in the others.

Tests.—Gallic Acid dissolves in Water, forming a solution which is acid in reaction towards blue Litmus paper, and which yields a bluish-black precipitate on the addition of a few drops of Ferric Chloride Test-solution. Solutions of pure Ferrous salts are unaffected by the addition of solution of Gallic Acid, but with Ferric salts precipitation takes place as above. The U.S.P. states that at about 200° C. (392° F.) it begins to melt.

The more generally occurring impurities are excess of Water, Tannic Acid, Sulphates and mineral matter. The crystalline acid should lose 10 per cent of its weight at a temperature of 100° C. (212° F.), indicating one molecule of Water of crystallisation. This statement is common to the B.P. and U.S.P. The absence of Tannic Acid is shown by the aqueous solution of the acid failing to give a precipitate with solutions of Isinglass or Albumen. The B.P. adds or Tartarated Antimony. The U.S.P. substitutes Solution of Gelatin for Solution of Isinglass, omits the Solution of Tartarated Antimony, but includes Test-solution of Starch. It also includes tests with Calcium Hydroxide Solution and with Sodium Hydroxide Solution, which are described below. It has been pointed out (P.J. '98, ii. 684, '99, i. 58) that what is ordinarily understood as 'Gallic Acid' gives a precipitate with Tartarated Antimony Solution, contrary to the official statement.

The absence of Sulphates is ensured by the usual tests, and the absence of mineral matter by the absence of ash when the acid is ignited at a low red heat.

Residue. When incinerated with free access of air, Gallic Acid should leave no residue, B.P.; U.S.P. states that at about 200° C. it begins to melt, and at a high temperature it is gradually decomposed, being consumed at a low red heat without leaving a residue.

Sodium Hydroxide Solution. If 6 drops of Sodium Hydroxide Test Solution be added to 5 c.c. of a saturated aqueous solution of Gallic Acid on a watch glass, the liquid will gradually acquire a deep green colour, which is changed to red or brownish-red by acids. Difference from and absence of Tannic Acid, U.S.P.

Calcium Hydroxide Solution.—When Calcium Hydroxide Test Solution is added to a cold saturated solution of Gallic Acid, a bluish white precipitate forms where the test solution is temporarily in excess and disappears on shaking. When the test solution has been added in excess the precipitate no longer dissolves, and the liquid acquires a tint that is blue by reflected and green by transmitted light and becomes pink on the addition of a large excess of Calcium Hydroxide Test Solution. Distinction from Tannic Acid, U.S.P.

Not Official.

GALLANOL (Gallic Acid Anhydride).—Colourless crystals, melting at 205° C. Insoluble in Water. Introduced as a substitute for Chrysophanic Acid in psoriasis. B.M.J. '03, ii. 99, '01, i. 12; ii. 44. In oozema.—M.A. '95, 296.

GALLOFORMIN.—A compound of Gallic Acid with Hexamethylenetetramine. Glistening needles, almost insoluble in cold Water. Used externally and internally as a disinfectant.

GALLOSOMOL (*Dibromogallic Acid*) (colourless needles or prisms, or as a white crystalline powder) Soluble 1 in about 8 of Water, readily soluble in Alcohol and Ether. Used internally as a substitute for the alkali Bromides in daily doses of 2 to 3 grammes (30 to 45 grains). Also in the form of a 1 to 2 p.c. solution as an injection in gonorrhoea.

Dose —8 to 16 grains = 0.52 to 1 gramme, three times a day.

Under the name of **Gallogen**, Ellagic Acid, the astringent principle of *Diri diri*, has been introduced as an astringent.

Not Official

ACIDUM GLYCEROPHOSPHORICUM.

See CALCI GLYCEROPHOSPHAS

Not Official

ACIDUM HYDRIODICUM.

This Acid is best prepared and kept in the form of a 20 p.c. solution (sp. gr. 1.17) by passing Hydrogen Sulphide gas through four parts of Water containing one part of Iodine. The action is rather slow at first, but becomes more rapid as more Iodine is dissolved by the Hydriodic Acid formed, till the absorption becomes very rapid. When the solution is colourless, the excess of Hydrogen Sulphide may be boiled off and the liquid filtered from separated Sulphur.

Though colourless when first made, it rapidly decomposes, even in diffused light, with liberation of Iodine, but may be readily decolorised by warming with a small proportion of Hypophosphorous Acid, 60 minims to 4 oz. is usually sufficient even for a highly coloured Acid.

Acidum Hydriodicum Dilutum (U.S.) — A solution containing not less than 10 p.c. w/w of absolute Hydriodic Acid.

SYRUPUS ACIDI HYDRIODICI — Colourless Hydriodic Acid (20 p.c.), $3\frac{1}{2}$ oz., Distilled Water, 8 oz., Simple Syrup, sufficient to make up the measure to 80 oz.

An acid syrupy liquid, colourless, or of a pale straw tint. Sp. gr. 1.300. Contains 1 p.c. of absolute Hydriodic Acid, HI. The U.S.P. preparation contains the same percentage of absolute Hydriodic Acid, but is of a much lower specific gravity (1.190 at 25° C (77° F)). A test is included for a limit of free Iodine, also a confirmatory test for Hydriodic Acid by means of Test solution of Silver Nitrate, and the amount of absolute Hydriodic Acid is determined by the addition of a definite volume of Tenth-normal Volumetric Solution of Silver Nitrate, a little diluted Nitric Acid, followed by the addition of Test solution of Ferric Ammonium Sulphate. The excess of Volumetric Silver Solution is determined by titration with Tenth-normal Volumetric Solution of Potassium Sulphocyanate.

Dose —20 to 40 minims = 1.2 to 2.4 c.c., well diluted.

Foreign Pharmacopœias. —Official in U.S. Not in the others.

Diluted Hydriodic Acid (10 p.c.), 10, Water, 30, Syrup, 60, all by weight to make 100 —U.S.P.

Diluted Hydriodic Acid (10 p.c.), 10, Water, 30, Syrup, to produce 100, all by weight —B.P.C.

ACIDUM HYDROBROMICUM DILUTUM.

DILUTED HYDROBROMIC ACID

FR., ACIDE BROMHYDRIQUE DILUÉ, GER., BROMWASSERSTOFFSAURE,
ITAL., ACIDO BROMIDRICO, SPAN., ACIDO BROMHÍDRICO OFICIAL.

A clear, colourless liquid, containing 10 p.c. by weight of Hydrogen Bromide, HBr, eq. 80.35.

It may be obtained by the decomposition of Potassium Bromide by concentrated Phosphoric Acid and distillation. Considerable trouble has been experienced (*P.J.* '00, 1 310, *V.B.P.* '00, 19) in obtaining pure Hydrobromic Acid by the use of red Phosphorus, owing to the Arsenic present in the latter. The method suggested in the reference consists of blowing Sulphur Dioxide into a mixture prepared by covering a quantity of pure Bromine with six times its volume of Water. On distilling the liquid the remaining Bromine goes over first, and is easily got rid of.

It should be preserved in stoppered glass bottles of a dark amber tint, and as far as possible protected from the light.

Medicinal Properties. Sedative and hypnotic, but not so reliable as the Bromides, though producing less depression. When continued sedative action is indicated, the acid can be used to supplement or replace the Bromide salts. It is stated to be less likely to produce acne.

Dr. Fothergill stated that it prevents headache after taking Quinine and Iron, and may be given with Quinine (which it readily dissolves) for nervous exhaustion.

It is said to prevent the after effects of Morphine if given with that drug.

Dose. --15 to 60 minims = 0.9 to 3.6 c.c.

Prescribing Notes.—*Larger doses may be given, 2 to 4 fl. drms, well diluted with Water, or Syrup and Water.*

60 minims = 8½ grams of Potassium Bromide in the quantity of Bromine.

Foreign Pharmacopœias.—Official in Dutch, sp gr 1.224, Fr., Span., Swiss and U.S., 10 p.c., sp gr 1.076 to 1.077, Ger. has 20 p.c., sp gr 1.208. Not in the others.

Tests.—Diluted Hydrobromic Acid has a specific gravity of 1.077, the *U.S.P.* states 1.076 at 25° C (77° F.), *P.G.* 1.208. A solution of the neutralised acid should give with Silver Nitrate Solution a yellowish curdy precipitate, insoluble in Nitric Acid, but soluble in Potassium Cyanide Solution, and soluble with difficulty in strong Ammonia Solution; Chlorine Solution causes a yellowish or reddish coloration due to the liberation of Bromine, which dissolves on shaking with a few drops of Chloroform or Carbon Bisulphide forming a reddish solution. It is officially required to indicate 9.98 p.c. of absolute Hydrobromic Acid as determined by titration with Volumetric Sodium Hydroxide Solution, or by precipitation with Volumetric Silver Nitrate Solution. The *P.G.* Volumetric test indicates an acid containing 25 p.c. by weight, and the *U.S.P.* test not less than 9.99 p.c. by weight of absolute Hydrobromic Acid. A comparison of the methods adopted by the *B.P.*, *U.S.P.* and *P.G.* appears below under the heading of Volumetric Determination.

The more generally occurring impurities are solid residue, Arsenic, Copper, Lead and Iron; Barium, Chlorides, Phosphates, Sulphates and Sulphites, impurities which are present in the materials used in the manufacture and escape removal during the purification of the acid. Mineral residue is readily detected by evaporation to dryness. Arsenic is the most important impurity, as Phosphorus

and Phosphoric Acid are both liable to contain this substance. The *USP* introduces a special test (the modified Gutzzeit's test) for Arsenic, and also requires that this should not respond to the time-limit test for heavy metals. The *BP* does not include a test for the latter. The *PG* includes a test for Iron with Potassium Ferrocyanide Solution, which is given in the small type below.

Copper and Lead, if present, may be detected by the test with Hydrogen Sulphide given in the small type below, and if the solution be made alkaline with Ammonia the test also affords an indication of the presence of Iion. Iodine, if present, may be detected by the test described under the heading of Chloroform. Barium, Chlorides, Phosphates, Sulphates and Sulphites may be detected by the tests in small type below under the respective headings of Potassium Sulphate Solution, Silver Nitrate Solution followed by Ammonium Carbonate Solution, Magnesium Sulphate Solution, Barium Nitrate or Chloride Solution.

Distillation.—The *USP* states that on distilling it, Water and a weak acid first pass over, when the temperature of 126°C (258.8°F) is reached an acid of 48 p.c. remains, which may be distilled unchanged.

Residue.—*BP* requires that it should yield no residue on evaporation to dryness. In this test the *USP* directs that after evaporation to dryness the temperature be brought to 110°C (230°F), when 10 c.c. of the acid should leave no appreciable residue.

Chloroform.—Hydrobromic Acid when shaken with Chloroform should not impart to the Chloroform a yellow colour, nor on the subsequent addition of a drop of Ferric Chloride Solution should a violet colour be produced. *PG*. The *USP* gives 10 c.c. of the Acid and 2 c.c. of Chloroform, and the *PG* uses equal volumes. Chloroform shaken with Hydrobromic Acid previously mixed with Chlorine Water is coloured a brownish-yellow. *PG*. The *USP* directs that Chlorine Water be diluted with an equal volume of Water and added drop by drop with agitation to 10 c.c. of Hydrobromic Acid and 2 c.c. of Chloroform, previously shaken together, when the Chloroform should be coloured orange, with no trace of violet, indicating the absence of Iodine.

Hydrogen Sulphide Solution.—Diluted with 5 volumes of Water, and nearly neutralised with Solution of Ammonia, Hydrobromic Acid should be unaffected by Hydrogen Sulphide Solution. *PG*. The *USP* requires that 10 c.c. of Diluted Hydrobromic Acid should not, without further acidulation, respond to the time-limit test for heavy metals.

Barium Nitrate or Chloride Solution.—10 c.c. of the Acid should not be rendered more than slightly cloudy by the addition of 1 c.c. of Barium Chloride T.S., indicating a limit of Sulphuric Acid. *USP*, *PG* nearly neutralises with Ammonia Solution, and uses Barium Nitrate Solution.

Potassium Sulphate Solution.—10 c.c. of the Acid should yield no turbidity with 1 c.c. T.S. of Potassium Sulphate, *USP*.

Magnesium Sulphate Solution.—1 c.c. of Acid with 1 c.c. of Nitric Acid, boiled, cooled, and then supersaturated with Solution of Ammonia should be unaffected by Magnesium Sulphate Solution even after standing for some time. *PG*.

Modified Gutzzeit's Test.—5 c.c. should not respond to the modified Gutzzeit's test for Arsenic.

Silver Nitrate Solution followed by Ammonium Carbonate Solution.—If a mixture of 0.5 c.c. Diluted Hydrobromic Acid, 10 c.c. of Water, 8 c.c. Silver Nitrate Test Solution, and 6 c.c. Ammonium Carbonate Test Solution, be digested for 10 minutes on a bath of boiling Water, then cooled and filtered, the filtrate when supersaturated with Nitric Acid should not become more than slightly opalescent, *USP*.

Potassium Ferrocyanide Solution. The *P.G.* requires that 10 c.c. of Hydrobromic Acid diluted with Water (1-10) should not immediately turn blue with 0.5 c.c. Potassium Ferrocyanide Solution.

Volumetric Determination.—The *B.P.*, *U.S.P.* and *P.G.* differ in their method of determining the amount of absolute Hydrobromic Acid. The *B.P.* employs both titration with Volumetric Sodium Hydroxide Solution and precipitation with Volumetric Silver Nitrate Solution, the *U.S.P.* employs direct titration with Tenth normal Volumetric Silver Nitrate, the *P.G.* titration with Normal Volumetric Potassium Hydroxide Solution. It is officially required that 4 grammes of diluted Hydrobromic Acid should neutralise 5 c.c. (1.08) of the Volumetric Sodium Hydroxide Solution, *B.P.*, 5 c.c. of the *P.G.* Acid should require 18.7 c.c. Normal Potassium Hydroxide Solution, *P.G.*. The *B.P.* requires that 4 grammes of the dilute Acid should be completely precipitated by 50 c.c. (49.8) of the Volumetric Solution of Silver Nitrate. The *P.G.* directs that 10 c.c. of a mixture of Hydrobromic Acid and Water (3 grammes in 100 c.c.) be exactly neutralised with Ammonia Solution and a drop of Potassium Chromate Solution added, when 9.3 c.c. at most of Dec normal Volumetric Silver Nitrate Solution are necessary to produce a permanent red colour, *P.G.*, the *U.S.P.* test is similar, 10 grammes of Acid being diluted to 100 c.c., then 8.01 c.c. of this solution exactly neutralised with diluted Ammonia Water (using Litmus Test solution as indicator), and 3 drops of Potassium Chromate Test solution added, should require not less than 10 c.c. of Tenth normal Silver Nitrate Volumetric Solution to impart a permanent red tint.

ACIDUM HYDROCHLORICUM.

HYDROCHLORIC ACID

Fr., ACIDE CHLORHYDRIQUE OFFICINAL; GER., SALZSÄURE, ITAL., ACIDO CLORIDRICO CONCENTRATO, SPAN., ACIDO CLORHIDRICO.

A colourless fuming liquid, containing 31.79 p.c. by weight of Hydrogen Chloride (HCl , eq. 36.49), possessing an irritating pungent odour, and even in dilute solutions an intensely acid taste.

It may be obtained by the decomposition of a Chloride, generally Sodium Chloride with Sulphuric Acid.

Acidum Hydrochloricum *B.P.* 1865 contained 32 p.c. of Hydrogen Chloride.

Medicinal Properties.—A powerful escharotic. When diluted it is given internally, *see* Acidum Hydrochloricum Dilutum.

Treatment of ulceration of the œsophagus and stomach due to swallowing strong Hydrochloric Acid.—*B.M.J.* '01, ii. 1168, '02, i. 511.

Treatment of lupus by thoroughly rubbing crude Hydrochloric Acid over the patch previously frozen by Ethyl Chloride.—*L.* '07, ii. 81.

Incompatibles.—Salts of Silver and Lead, Tartar Emetic, Alkalies and their Carbonates.

Official Preparations.—Acidum Hydrochloricum Dilutum.* Used in the preparation of Acidum Nitro-hydrochloricum Dilutum, Apomorphinæ Hydrochloridum, Cocainæ Hydrochloridum, Extractum Cinchonæ Liquidum, Glycerinum Populæ, Liquor Arsenici Hydrochloricus, Liquor Ferri Perchloridi Fortis, Liquor Zinci Chloridi, and Podophylli Resina.

Antidotes.—In cases of poisoning by Hydrochloric Acid, the antidotes are Chalk, Magnesia, Potassium Bicarbonate, with Whites of Egg, Carron Oil, or Soap-suds, followed by enemata of Beef Tea and Brandy (with Tincture of Opium) to prevent collapse; and emollient drinks.

Foreign Pharmacopœias.—Official in Austr., 25 p.c., *sp. gr.* 1.194; Belg., *sp. gr.* 1.186, Port. and Spain, *sp. gr.* 1.180; Dutch and Swiss, 25 p.c., *sp. gr.*

1 126, Fr, 83 65 p.c., sp. gr 1.171, Jap, 80 p.c., sp. gr 1 15, Mex, 1 17, Dan, Norw and Swed, 25 p.c., sp. gr 1 127, Ger, Hung and Russ, 25 p.c., sp. gr 1 124, Ital, 35 39 p.c., sp. gr 1 18, U.S., 31 9 p.c., sp. gr 1 158 at 25° C (77° F) An **Acidum Hydrochloricum Crudum** is included in the Fr, Ital, Russ and Swed.

The Crude Acid made with Pyrites Vitriol is generally yellow, and contains considerable traces of Iron and Arsenic

Tests.—Hydrochloric Acid has a specific gravity of 1 160, the *USP* gives about 1 158 at 25° C (77° F), the *PG* 1 124 The diluted or neutralised solutions afford, when treated with Silver Nitrate Solution, a curdy white precipitate, insoluble in dilute Nitric Acid, but readily soluble in Ammonia Solution Another characteristic test for Hydrochloric Acid, which in the *PG* and *USP* is performed upon the pure undiluted or unneutralised acid, but in the *BP* appears amongst the miscellaneous collection of tests suitable for application to the neutralised acid, is that when warmed with Manganese Oxide, Chlorine gas is evolved, which may be recognised by its colour, odour, and bleaching action upon moistened Litmus paper, and by liberating Iodine when brought into contact with Potassium Iodide Solution The acid is officially required to contain 31.49 p.c. of absolute Hydrochloric Acid as indicated by titration with Volumetric Solution of Sodium Hydroxide, the *USP* test indicates 31.9 p.c. by weight of absolute Acid, and the *PG* 25 p.c. by weight The processes are compared below under the heading of Volumetric Determination

The more generally occurring impurities are mineral residue, Arsenic, Lead, Copper, Iron, Aluminium, free Chlorine, Bromine, Iodine, Sulphates and Sulphurous Acid The most important of these are Arsenic, Iron, and free Chlorine Mineral matter is readily detected by the residue left on evaporation The *BP* characteristically groups Arsenic amongst the general list of impurities, and employs the tests mentioned in the Appendix for its detection The *USP* adopts the modified Gutzeit's test, and the *PG* the test with Stannous Chloride Solution The *PG* includes a specific test for Iron with Potassium Ferrocyanide Solution, see below The three Pharmacopœias differ in their manner of testing for free Chlorine All three use the diluted acid, the *PG* in addition partially neutralising the liquid, the *BP* employs Potassium Iodide and Starch Solution as a reagent, the *PG* Zinc Iodide and Starch Solution, the *USP* uses Potassium Iodide Solution, but shakes with Chloroform, and notes the absence of a violet coloration in the chloroformic layer in preference to the Starch test A 1 to 20 dilution of the Acid almost neutralised with Ammonia Solution should not be altered by the addition of Hydrogen Sulphide Solution, indicating the absence of Lead and Copper The *USP* requires that the 1 in 20 aqueous dilution should not respond to the time limit test for heavy metals, the *PG*, that the 1 to 5 dilution, when almost neutralised with Ammonia Solution, shall not be altered by Hydrogen Sulphide Solution Bromine and Iodine, Sulphates and Sulphites may be detected by the respective tests under the headings of Chlorine Water, and Barium Chloride or Nitrate Solution

A standard of 5 parts per million for Arsenic is suggested (*C D* '08, i 795) as sufficient for a pharmaceutical acid, and 10 parts per million for Lead

Stannous Chloride.—A mixture of 1 c.c. of Acid and 3 c.c. Stannous Chloride Solution should not assume a dark colour in the course of an hour, *P G*

Modified Gutzzeit's Test.—5 c.c. of diluted Acid (1:10) should not respond to the modified Gutzzeit's test for Arsenic

Potassium or Zinc Iodide Solution.—When largely diluted with Water it should yield no blue coloration on the addition of Potassium Iodide Solution and Starch Mucilage, *B P*, the *P G* directs the Acid to be diluted with 5 volumes of Water, and nearly neutralised with Solution of Ammonia, and requires that this solution should not immediately turn blue with a Solution of Zinc Iodide and Starch. The *U S P* dilutes 1 c.c. of Acid with 5 c.c. of Water, then on the addition of 1 c.c. of Potassium Iodide T.S. and 1 c.c. of Chloroform, and the mixture agitated, the Chloroform should be free from any violet coloration

Chlorine Water. If Chlorine Water diluted with an equal quantity of Water be added cautiously, drop by drop with constant agitation, to Hydrochloric Acid diluted with an equal volume of Water, and 1 c.c. of Chloroform added, the Chloroform should be free from any yellow, orange, or violet colour, *U S P*

Barium Chloride or Nitrate Solution.—Hydrochloric Acid diluted 1 to 5 with Water and nearly neutralised with Ammonia Solution should be unaffected within 5 minutes by Barium Nitrate Solution, *P G*. The *U S P* uses Barium Chloride T.S., a few drops added to 1 c.c. of the Acid previously diluted with 5 c.c. of Water, when no turbidity or precipitate should be produced within 1 hour, nor should the addition of a few drops of Tenth-normal Iodine V.S. to the mixture produce any turbidity.

Potassium Ferrocyanide Solution.—10 c.c. Hydrochloric Acid diluted with Water (1-10) should not immediately turn blue on the addition of 0.5 c.c. Potassium Ferrocyanide Solution, *P G*

Volumetric Determination (by neutralisation).—The *B P* requires that 1 gramme diluted with Water should neutralise 8.7 c.c. of the Sodium Hydroxide Volumetric Solution, precipitation with Volumetric Silver Nitrate Solution is also adopted as a means of determining the amount of absolute Acid present, 8.7 c.c. of Volumetric Silver Nitrate Solution being required to precipitate 0.1 of a gramme of the acid. The *P G* uses Normal Potassium Hydroxide Solution, 38.5 c.c. of this solution being necessary for 5 c.c. of Acid. The *U S P* directs that 8 c.c. of Acid be accurately weighed, diluted with 5 c.c. of Water and titrated with Normal Potassium Hydroxide Solution, using Methyl Orange Test-solution as indicator. The number of c.c. required, multiplied by 3.618 and divided by the weight of Acid taken, represents the percentage of absolute Hydrochloric Acid in the sample taken

Preparation.

ACIDUM HYDROCHLORICUM DILUTUM. DILUTED HYDROCHLORIC ACID.

Dilute 6 of Hydrochloric Acid with Distilled Water to make 20.

34 minims contain about 1 minim of Strong Acid

It is a clear, colourless and odourless liquid, possessing a strong acid taste and acid reaction to Litmus

Medicinal Properties.—Stimulant, sialagogue, stomachic tonic, cholagogue. Externally and diluted it is refrigerant. Given about two hours after meals in dyspepsia due to deficient secretion of Hydrochloric Acid, given before meals in cases of acid eructation and heartburn, to prevent excessive secretion of acid, used also in gargles; given internally also to diminish night sweating.

10 to 20 drops of a 38 p.c. solution of pure Hydrochloric Acid in a little simple Syrup at the beginning of a meal in the treatment of chronic diarrhoea — *B M J E '02*, 11 7

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

Prescribing Notes.—Usually given with aromatic or bitter infusions, for children, $1\frac{1}{2}$ to 2 minims, 1 drm in 8 oz of Infusion of Roses or Decoction of Cinchona as a gargle for ulcerated sore throat

Official Preparations.—Used in the preparation of Extractum Ergotæ, Injectio Apomorphinæ Hypodermica and Liquor Morphinæ Hydrochloridi

Foreign Pharmacopœias—Austr., 12.5 p.c., sp. gr. 1.061, Belg., sp. gr. 1.037, Fr., Hung., Swiss and U.S., 10 p.c., about sp. gr. 1.049, Dutch, sp. gr. 1.037, Jap., 10 p.c., sp. gr. 1.050, Dan., Norw. and Swed., 10 p.c., sp. gr. 1.050 to 1.052, Ger., 12.5 p.c., sp. gr. 1.061, Ital., 8.07 p.c., sp. gr. 1.036, Russ., 8.2 p.c., sp. gr. 1.040, Mex., Acid 1, Water 3 Not in the others

Tests—Diluted Hydrochloric Acid has a specific gravity of about 1.052. When its neutralised or diluted aqueous solution is mixed with Silver Nitrate Solution, a curdy white precipitate insoluble in Nitric Acid, but readily soluble in Ammonia Solution and in Potassium Cyanide Solution is produced.

As the diluted Acid is directed to be prepared with the official Hydrochloric Acid, it is required to be free from the impurities mentioned under the concentrated Acid. It is officially required to contain 10.49 p.c. by weight of absolute Hydrochloric Acid, as indicated by titration with Normal Volumetric Sodium Hydroxide Solution

ACIDUM HYDROCYANICUM DILUTUM.

DILUTED HYDROCYANIC ACID

HCN, eq. 26.85

FR., ACIDE CYANHYDRIQUE DISSOUS, GER., CYANWASSERSTOFFSAURE,
ITAL., ACIDO CIANIDRICO, SPAN., ACIDO CIANHIDRICO MEDICINAL

A clear, colourless liquid, possessing a characteristic odour somewhat resembling bitter almonds. It is officially required to contain 2 p.c. by weight of Hydrogen Cyanide, and may be prepared by the interaction of Potassium Ferrocyanide and Sulphuric Acid, or when only a small quantity is required occasionally, it may be convenient to prepare it extemporaneously from dry Silver Cyanide, as in the *U.S.P.*—Silver Cyanide, 6 parts, Diluted Hydrochloric Acid (*B.P.*), 15.54 fluid parts; Distilled Water, 44.1 parts. Shake for a short time and filter. The product should contain 2 p.c. w/w of Hydrogen Cyanide.

It should be kept in dark amber-tinted, well stoppered glass bottles, in a cool and dark place, the bottles being maintained in an inverted position.

Medicinal Properties.—As this Acid is a dangerous poison, it should never be prescribed undiluted. Moreover, a diluted solution retains its strength better than a strong one.

It is sedative, antispasmodic, allays vomiting, is useful in gastrodynia, in visceral neuralgias, in dyspeptic palpitations, but

chiefly valuable in the dry resultless cough of asthma, phthisis and whooping-cough, and prevents the vomiting brought on by food in phthisis. Used externally to allay itching in urticaria, lichen, etc., if the skin be unbroken, as a **lotion**, 2 drin to 8 oz of Rose Water and Glycerin; as an **ointment**, from $\frac{1}{2}$ to 1 drin to each oz of Zinc ointment.

The **vapour** is sometimes applied to the eye, but it is more generally used as a sedative inhalation in the cough of laryngeal phthisis and in some spasmodic affections.

Dose.— 2 to 6 minims = 0.12 to 0.36 c c

Prescribing Notes.— Given in Almond Emulsion for cough, and with Sodium Bicarbonate, Bismuth Carbonate and Peppermint Water for dyspnoea.

As it is susceptible to the action of light and air, and is very volatile, it is the practice to keep it in amber-coloured bottles, stopper down wards.

Incompatibles. Silver, Copper, and Iron salts, and Mercuric Oxide.

Official Preparations.— Used in the preparation of *Imctura Chloroformi* et *Morphine Composita*.

Not Official.— *Acidum Hydrocyanicum* (Schell). Brompton Cough Mixture, *Mistura Acidi Hydrocyanici Composita*.

Antidotes.— In cases of poisoning, the antidotes are fresh air and artificial respiration, with cold affusion, the recent precipitate obtained by swallowing 10 grains of Ferrous Sulphate with 1 fl drin of Tincture of Ferric Chloride in 1 oz of Water, followed by 20 grains of Potassium Carbonate dissolved in 1 oz. of Water, this will render insoluble 110 minims of *H.P.* Acid Stimulants, Ammonia and Brandy, Hypodermic injection of Atropine, $\frac{1}{10}$ grain.

Suggested that in mines and places where Cyanide is used the following antidote should be kept ready: (1) 1 oz of a 24 p.c. solution of Ferrous Sulphate, (2) 1 oz. of a 5 p.c. solution of Potassium Hydrate, (3) 80 grains of powdered Magnesium Oxide to be added to the above in half a pint of Water — *J. OI.*, n. 197.

Injection of a 3 p.c. solution of Hydrogen Peroxide simultaneously recommended in cases of poisoning by fumes of Hydrocyanic Acid — *J.C.S.*, '01, *Abs.* n. 535.

Foreign Pharmacopœias. Official in Belg., 2.0 p.c., Dutch, 2 p.c.; Fr., 2 p.c., Jap. 2 p.c., Norw. 2 p.c., Port., strength not given, Mex. (Acido cianhidrico medicinal); U.S., 2 p.c., Span., 2 p.c. Not in the others. See also *Aqua Amygdale Amare*.

The *Brussels Conference* adopted a strength of 2 p.c. for *Acidum Hydrocyanicum Dilutum*.

Tests.— Hydrocyanic Acid has a specific gravity of about 0.997; the *U.S.P.* does not give a specific gravity; the acid is not included in the *P.G.* When Silver Nitrate Solution is added to its neutralised solution there is produced a white curdy precipitate, soluble in Potassium Cyanide Solution, in Ammonia Solution, and in Nitric Acid. A blue precipitate is produced when a mixture of Ferrous and Ferric salts in solution is added to the neutralised acid, followed by the addition of Sodium Hydroxide Solution and then an excess of Hydrochloric Acid.

The Acid is officially required to contain 1.98 p.c. by weight of absolute Hydrocyanic Acid as indicated by titration with Volumetric Silver Nitrate Solution after the liquid has been rendered alkaline by the addition of Sodium Hydroxide Solution, 1 gramme of the Acid should require 3.7 c.c. of Deci-normal Volumetric Silver Nitrate

Solution This process of titration is not altogether satisfactory, and, moreover, estimates any Chloride present. A very useful method for determining the strength of Diluted Hydrocyanic Acid is —Place 10 c.c. of Ammonia Solution in a beaker, add 40 c.c. of Water and 0.2 grammes of Potassium Iodide and 5 c.c. of the Acid to be tested, titrate with Volumetric Silver Nitrate Solution, of which 18.7 c.c. will be required for a 2 p.c. Acid. The presence of Hydrochloric Acid (a trace of which is understood to be purposely added by manufacturers to retard decomposition) will not affect the results of the test, and the end reaction is very definite.

The more generally occurring impurities are mineral residue, Sulphates and Chlorides. The mineral residue is readily determined by evaporating the fluid to dryness.

When diluted with Water and slightly acidified with Hydrochloric Acid Solution it should not yield a distinct turbidity on the addition of Barium Chloride Solution, and when acidified with Nitric Acid should not yield a distinct turbidity on the addition of Silver Nitrate Test-solution, indicating the absence of more than traces of Chlorides and Sulphates.

Volumetric Determination—A weighed quantity of 5 grammes of the Acid is diluted to 50 c.c. with Water, 26.9 c.c. (26.84) of this solution, after the addition of 5 c.c. of Ammonia Water and 3 drops of Potassium Iodide T.S., should require for the production of a slight permanent precipitate the addition of not less than 10 c.c. Dec. normal Volumetric Silver Nitrate Solution, *U.S.P.*

The *B.P.* and *U.S.P.* Acids contain not less than 2 p.c. by weight of Absolute Hydrocyanic Acid.

Not Official

BROMPTON COUGH MIXTURE—Diluted Hydrocyanic Acid, 2½ minims, Solution of Morphine Hydrochloride, 7½ minims, Syrup of Tolu, 40 minims, Acid Infusion of Roses, to make 4 fl. oz.—*Pharm. Form.*

This has been incorporated in the *B.P.C.* under the title **Mistura Acidi Hydrocyanici Composita** with the *syn.* Brompton Hospital Mixture.

ACIDUM HYDROCYANICUM (SCHEELE) *B.P.C. Formulary 1901*—A colourless liquid. Sp. gr. 0.991. It should contain 4 p.c. HCN, as volumetrically determined by Volumetric Silver Nitrate Solution, it should give no precipitate with Barium Chloride Solution, but with Silver Nitrate a white precipitate entirely soluble in boiling concentrated Nitric Acid.

Dose—1 to 3 minims = 0.06 to 0.24 c.c.

This has been incorporated in the *B.P.C.* under the title **Acidum Hydrocyanicum Fortius**.

It is known that the weaker strengths of Hydrocyanic Acid keep better than the stronger, and the only practical use for a double strength Acid is to poison dogs or cats.

Not Official.

ACIDUM HYDROFLUORICUM.

A colourless liquid, usually redistilled, containing about 80 p.c. of Hydrofluoric Acid gas, owing to its action on glass it is usually stored in gutta-percha bottles.

It is strongly corrosive. Great caution must be used in handling this Acid, as contact with the liquid or gas may result in sores difficult to heal, or permanent destruction of tissue, no pain is felt until the injury is beyond remedy. It also gives off a pungent irritating vapour.

In experiments made to determine the most suitable indicator for the titration of Hydrofluoric Acid, Phenolphthalein Solution answered well with Potassium or Sodium Hydroxide. Rosolic Acid Solution was equally useful and had the additional advantage of being capable of use with Ammonia. Cochineal and Brazil Wood Solutions answered fairly well, but Methyl Orange Solution was useless. With Litmus Solution the colour change is somewhat complicated — *P. J.* (4) xxx 701.

ACIDUM HYDROFLUORICUM DILUTUM.—Dilute the 30 p.c. Acid so as to contain 0.2 p.c., and preserve in gutta serena bottles — *B. P. C. Formulary* 1901.

Dose.—5 to 20 minims = 0.3 to 1.3 c.c.

This has been incorporated in the *B. P. C.* giving the dose 5 to 15 minims.

AMMONII FLUORIDUM.—Colourless crystals soluble in Water. Given in hypertrophy of the spleen and in goitre as a $\frac{1}{2}$ p.c. solution, in doses of 5 to 20 minims = 0.3 to 1.3 c.c.

FERRI FLUORIDUM.—A mixture of Ferric and Ferrous Fluoride. A purplish-grey powder, insoluble in Water.

Under the proprietary titles of *Antitussin* and *Fluorheumum*, bodies containing Fluorine have been introduced, the former as an application in whooping cough, the latter for rheumatism — *P. J.* '99, n. 11; '00, n. 775.

Fluoroform (CHF_3) is analogous in composition to Chloroform (CHCl_3), but contains the halogen Chlorine replaced by Fluorine (CHF_3). The pure product is at ordinary temperatures a gas, and is prepared by the action of Silver Fluoride on Iodoform, the gas being subsequently purified from Carbonic Oxide by passing through a solution of Cuprous Chloride. This gaseous Fluoroform was tested pharmacologically and proved to be very similar in action to Chloroform. The commercial article is a 2.8 p.c. aqueous solution of the gas, and it is in this form that it is generally employed. It possesses very little odour or taste, and is comparatively harmless even in large doses. It has been mostly used in cases of phthisis, lupus, and tuberculous diseases of the joints. It is administered in doses of 1 to 3 teaspoonfuls taken in Water 4 or 5 times a day.

Sodium Fluoride has been introduced in the treatment of tuberculosis and stated to possess distinct antiseptic properties. Toxic effects of same — *P. J.* '99, n. 235. The Fluorides have been used as preservatives of foods.

Solutions of **Sodium Silicofluoride** and the **Saluferbath**, which is a mixture of various silicofluorides, have been introduced as antiseptics — *B. M. J. E.* '08, i. 712.

Not Official.

ACIDUM HYPOPHOSPHOROSUM.

H_2PO_2 , eq. 65.56.

A clear, colourless liquid, possessing an acid taste and an acid reaction to Litmus. It contains 30 p.c. of absolute Hypophosphorous Acid. It should be kept in dark amber-tinted, well-stoppered bottles and in a cool place.

Dose.—2 to 5 minims = 0.12 to 0.3 c.c.

A good preservative for preparations otherwise liable to change by oxidation.

Used principally in the manufacture of the Solution and Syrup of Iron Hypophosphite, etc.

Dissolve 8 oz. of Barium Hypophosphite (containing not less than 95 p.c. $\text{Ba}_2(\text{PH}_2\text{O}_2)_2 \cdot \text{H}_2\text{O}$) in 86 fl. oz. of hot Distilled Water. Add slowly to the solution 17 fl. oz. of Diluted Sulphuric Acid, after which continue the addition, drop by drop, until no further turbidity is produced. Set aside in a warm place, and pass the clear liquid through a filter. Wash the precipitate by decantation with successive portions of hot Distilled Water, until the washings have no longer an acid reaction. Filter, unite the filtrates, and evaporate the liquid on a water-bath to the prescribed density. The product will weigh about 11½ oz.

The process is better than the treatment of Calcium Hypophosphite with Oxalic Acid. But still a pure Hypophosphorous Acid is a commercial desideratum.

Tyrer compares the Barium and Calcium methods, and decides in favour of Barium.—*P J* '96, ii 94

Used in the manufacture of the Solution and Syrup of Iron Hypophosphite, etc

Foreign Pharmacopœias.—Official in U S contains 80 p c by weight, sp gr 1.180 at 25° C (77° F)

Tests.—Sp gr 1.1867 Its strength as determined by Volumetric Sodium Hydroxide Solution corresponds to 80 p c of absolute Hypophosphorous Acid. Its aqueous solution is not precipitated by Diluted Sulphuric Acid, nor by an excess of Ammonia Solution, nor by Ammonium Oxalate Solution after neutralisation, and gives not more than a faint opalescence with Barium Chloride Solution. If Magnesium Ammonio-sulphate Solution be added after an excess of Ammonia Solution, no precipitate is produced. Calcium Chloride Solution added to a neutralised solution yields no precipitate.

Heated with excess of Mercuric Chloride Solution and a little Hydrochloric Acid to 100° C (212° F), Calomel is precipitated, from the weight of which the percentage of Hypophosphorous Acid may be calculated.

As the reaction follows the equation $H_2PO_3 + 4HgCl_2 + 2H_2O = H_2PO_4 + 4HgCl + 4HCl$, 100 parts of Calomel produced are equivalent to 7 parts of Anhydrous Acid.

Acidum Hypophosphorosum Dilutum.—Hypophosphorous Acid, 20, Distilled Water, 40, both by weight.—*USP*

Sp. gr 1.042 at 25° C (77° F) contains 10 p c by weight of absolute Hypophosphorous Acid.

Hypophosphorous Acid, 83, Distilled Water, *q s* to produce 100.—*B P C*

This is obviously intended to yield the same result as the *USP* given above, but in that case the quantities should be by weight and not by volume.

ACIDUM LACTICUM.

LACTIC ACID

FR, ACIDE LACTIQUE, GLR, MILCHSAURE, ITAL, ACIDO LATTICO;

SPAN, ACIDO LACTICO

A colourless and odourless syrupy hygroscopic liquid, possessing a purely acid taste. It should contain 75 p c of Hydrogen Lactate, $HC_2H_3O_3$, eq 89.37

It is produced by the fermentation of Lactose, and is extracted in the form of Zinc Lactate, the latter salt being subsequently decomposed. It is also obtainable by various synthetical processes.

Solubility—It is miscible in all proportions with Water, Alcohol (90 p c), and Ether. It dissolves, but is not dissolved by, Chloroform.

Medicinal Properties—It is used as a 'swab' in diphtheria, a solution (50 to 75 p c) has been used successfully for pharyngeal and laryngeal tubercle, and for lupus after scraping.

50 p c solution applied to corneal ulcers.—*L* '95, i 1452

A 2 p c solution is recommended in the treatment of laryngeal papillomata.—*B M J* '04, ii 1224

Official Preparation Syrupus Calcii Lactophosphatis

Not Official.—Calcii Lactas, Ferri Lactas, Plumbi Lactas, Sodii L., Zinci L., Bismuthi L., Acidum Lacticum Dilutum, Syr. Calcii Lactophosphatis & Ferro

Foreign Pharmacopœias.—Official in Fr, sp gr 1.24, Port and Span, sp gr 1.215, Austr., Belg., Dan., Dutch, Ger., Ital., Jap., Norw., Russ., Swed. and Swiss, sp gr. 1.21 to 1.22, U S, sp gr. 1.206 at 25° C (77° F), Mex., sp gr. 1.315

Tests.—Lactic Acid has a specific gravity of 1.210, the *U.S.P.* gives 1.206 at 25° C (77° F), the *P.G.* 1.210 to 1.220. It vaporises at a temperature above 148.9° C (300° F), giving off inflammable vapours at a temperature of 176.7° C. (350° F), which on ignition burn with a blue flame. The readily recognised odour of Aldehyde is evolved when the Acid is warmed with Potassium Permanganate. The *B.P.* and *U.S.P.* employ for this test solid Potassium Permanganate, but the *U.S.P.* warms it with a mixture of equal volumes of the Acid, Potassium Permanganate, and Sulphuric Acid, whereas *B.P.* uses Lactic Acid and Potassium Permanganate; the *P.G.* uses a Potassium Permanganate Solution and Lactic Acid. It is officially required to contain 74.18 p.c. by weight of absolute Lactic Acid, as determined by titration with Volumetric Sodium Hydroxide Solution. It will therefore be noticed that the *B.P.* statement, 'a liquid containing 75 p.c. of Hydrogen Lactate,' is at variance with the volumetric determination. In the case of the Acid the *B.P.* has not distinctly specified 'per cent. by weight,' though this is clearly intended. Using the atomic weights official in the *B.P.* 1885 in calculating the result of the volumetric test, the official figure would indicate 74.7 p.c. by weight of absolute Acid. It seems, therefore, as if this were another instance in which the *B.P.*, after adopting new atomic weights, have omitted to bring their monographs into accordance with them. The test would read better, 'each gramme should require for neutralisation 8.4 c.c. of the Volumetric Sodium Hydroxide Solution.' The *U.S.P.* acid is required to contain not less than 75 p.c. by weight, the *P.G.* about 75 p.c. by weight of absolute Acid.

The more generally occurring impurities are heavy metals, *e.g.*, Arsenic, Copper, Lead and Iron, Sacco-lactic, Malic, and Sulphuric Acids, Glycerin, Cane, Grape or Milk Sugar, Fatty Acids, organic impurities, Calcium Phosphate, Gum, Mannite, mineral residue, Chlorides, Citrates, Oxalates, Phosphates, Sulphates, or Tartrates.

Heavy metals are grouped collectively in the *B.P.*, in the *U.S.P.* they are covered by the time-limit test, and by the Hydrogen Sulphide test described below. *B.P.* and *U.S.P.* test for Sacco-lactic Acid with Copper Sulphate Solution, but no similar test is included in the *P.G.*

Tests for Malic and Sulphuric Acids are not included in the *U.S.P.* or *P.G.* The *B.P.* uses Lead Subacetate Solution as a reagent requiring that no precipitate should be produced.

The tests adopted by the *B.P.* and *U.S.P.* in examining for Glycerin are essentially the same, and consist in converting the acid into a Zinc salt, drying and extracting with Absolute Alcohol, the *P.G.* does not include a test for Glycerin.

Cane, Grape and Milk Sugars are readily detected by their reducing action on Potassio-cupric Tartrate Solution, and the method of carrying out the test is essentially the same in the *B.P.* and *U.S.P.*, the latter Pharmacopœia definitely stating the relative quantities to be employed. The Fehling test does not appear in the *P.G.*

Foreign fatty acids are tested for in an almost identical manner by the three Pharmacopœias by warming the acid and observing

the odour, as are also organic impurities by the Sulphuric Acid test, the *USP* and *PG* allowing a limit of time (15 minutes) within which no darkening in colour shall take place, and the *USP* in addition maintaining the temperature of the mixed Sulphuric and Lactic Acids at or below 15° C (59° F)

Advantage is taken of the insolubility of Calcium Phosphate, Gum, Mannite and Sugar in Ether to utilise the latter substance as a test for them in the Acid. The test appears in the *BP* and *PG*, but not in the *USP*. The three Pharmacopœias differ in the amount of mineral residue permitted. The *BP* allows 0.5 p.c., the *PG* no weighable residue from 0.5 gramme, and the *USP*, 1.0 p.c.

Chlorides, Citrates, Oxalates, Phosphates, Sulphates or Tartrates are readily detected by the tests given in the small type below under the respective headings of Silver Nitrate, Lime Water, and Barium Chloride

Warming.—When gently warmed the Acid should not evolve the odour of rancid fat. Indicating the absence of Butyric and other fatty acids, *BP*, *PG* and *USP*

Potassium Permanganate—On warming with Potassium Permanganate (Potassium Permanganate Solution, *PG*), Lactic Acid evolves the odour of Aldehyde. *BP* and *PG*, the *USP* test directs equal volumes of Lactic and Sulphuric Acids and Potassium Permanganate

Sulphuric Acid—If Lactic Acid be carefully poured on an equal volume of Sulphuric Acid in a clean test tube (previously rinsed out with Sulphuric Acid, *PG*), no darkening in colour should occur (*BP*) within 15 minutes, indicating the absence of more than traces of organic impurities, *PG* and *USP*. *USP* also directs that the temperature be maintained at or below 15° C (59° F)

Copper Sulphate Solution—Diluted with Water (1-10 *PG* and *USP*) Lactic Acid should be unaffected by Copper Sulphate T.S., indicating the absence of Sarcosine, Lactic Acid, *BP* and *USP*

Potassium cupric Tartrate Solution—Diluted with Water the Acid should give no precipitate, or only the slightest, even after prolonged boiling with Potassium-cupric Tartrate Solution, *BP*, the *USP* orders a few drops of the Acid to be added to 10 cc of the hot Alkaline Cupric Tartrate Volumetric Solution, indicating the absence of Grape, Cane and Milk Sugars

Zinc Carbonate—If Lactic Acid be heated with an excess of Zinc Carbonate and evaporated to dryness (dried at 100° C (212° F), *USP*), and if this mixture be exhausted with Absolute Alcohol, and the alcoholic liquid evaporated, no sweet residue should be left, indicating the absence of Glycerin, *BP* and *USP*

Ether.—If Lactic Acid be added drop by drop to twice its volume of Ether, the mixture should not show any permanent or transient turbidity, indicating absence of Gum, Sugar, Mannite, Calcium Phosphate, *BP* and *PG*

Volumetric Determination—One gramme neutralises 8.8 cc. of the Volumetric Sodium Hydroxide Solution, *BP*. If 5 grammes be diluted to 50 cc with Water, then 41.7 cc of this solution should require for complete neutralisation at boiling temperature not less than 37.5 cc. of Potassium Hydroxide Solution (each cc = 2 p.c. absolute Lactic Acid), Phenolphthalein T.S. being used as indicator, *USP*

Barium Chloride or Nitrate Solution.—10 cc. of an aqueous solution of a strength of 1 in 20 should be unaffected by 1 cc of Barium Chloride Test-solution, *USP*, or by Barium Nitrate Solution, *PG*, indicating the absence of Sulphates.

Hydrogen Sulphide Solution.—An aqueous solution of a strength 1 in 10 shall not be altered by the addition of Hydrogen Sulphide Solution, *P.G.*, nor should an aqueous solution of similar strength respond to the tume limit test for heavy metals, *U.S.P.*, indicating the absence of Arsenic, Copper, Lead and Iron.

Silver Nitrate Solution.—An aqueous solution of a strength of 1 in 10 shall not be altered by the addition of Silver Nitrate Solution, *P.G.*, 10 c.c. of an aqueous 1 in 100 solution should not be rendered opalescent by 1 c.c. of Silver Nitrate Solution, *U.S.P.*

Ammonium Oxalate Solution.—An aqueous 1 in 10 solution shall not be altered by the addition of Ammonium Oxalate Solution, indicating the absence of Calcium, *P.G.*

Lime Water.—An aqueous 1 in 10 solution shall not be altered by the addition of excess of Lime Water even on heating, *P.G.*

Preparation.

SYRUPUS CALCII LACTOPHOSPHATIS. SYRUP OF CALCIUM LACTOPHOSPHATE.

Precipitated Calcium Carbonate, 2½ oz.; Concentrated Phosphoric Acid, 4 fl. oz. and 262 minims, Lactic Acid, 6 fl. oz., Refined Sugar, 70 oz.; Orange Flower Water of commerce, undiluted, 2½ fl. oz.; Distilled Water, sufficient to produce 100 fl. oz. of Syrup.

The Lactic Acid is diluted with four times its volume of Distilled Water, and the Calcium Carbonate is added in small portions. As soon as the Carbonate is completely in solution the concentrated Phosphoric Acid is added and trituration continued until the precipitate at first formed redissolves. The solution is diluted with some Distilled Water, the undiluted Orange Flower Water added, and the mixture filtered. The Refined Sugar is dissolved without the aid of heat in the filtrate, strained, and sufficient Distilled Water added to produce 100 fl. oz. of Syrup.

Dose.—30 to 60 minims = 1.8 to 3.6 c.c.

The occasional change of colour in this syrup is stated to be due to inversion of the Sugar by the Acid—*P.J.* '99, 11 221, *A.J.P.* '98, 589.

Foreign Pharmacopœias.—Official in Belg., contains 1.5 p.c. Bi Calcium Phosphate, Mex., and Spain contain 1.25 p.c. by weight of Bi Calcium Phosphate; Swiss and U.S. from Calcium Carbonate, *see* below.

Not Official.

ACIDUM LACTICUM DILUTUM.—Lactic Acid, 3 fl. oz.; Distilled Water, *q.s.* to produce 20—*B.P.* 1885.

This has been incorporated in the *B.P.C.*

SYRUPUS CALCII LACTOPHOSPHATIS.—Precipitated Calcium Carbonate, 2.5, Lactic Acid, 6; Phosphoric Acid, 3.6, Orange Flower Water, 5, Sugar, 72.5, Water, *q.s.* to make 100.—*U.S.P.*

Calcium Carbonate, 1, Lactic Acid, 2.4, Diluted Phosphoric Acid, 2, Simple Syrup, 80, Water, *q.s.* to make 100. All by weight—*Swiss*.

SYRUPUS CALCII LACTOPHOSPHATIS C. FERRO.—Ferrous Lactate, 8.5, Potassium Citrate, 8.5, Water, 62.5, Syrup of Calcium Lactophosphate (*U.S.P.*), *q.s.* to make 1000—*U.S.N.F.* 1906.

The *B.P.C.* has incorporated the formula of the *U.S.N.F.* 1906, but employs *B.P.* Syrup of Calcium Lactophosphate in place of the *U.S.P.* Syrup as follows.—*B.P.* Ferrous Lactate, 0.85, Potassium Citrate, 0.85; Distilled Water, 6, Syrup of Calcium Lactophosphate, *q.s.* to make 100.

Other Not Official salts of Lactic Acid are:—**Plumbi Lactas**, a heavy white crystalline powder, soluble in Water. **Sodium Lactate**, a colourless or

light yellow liquid of a syrupy consistency and mild salty taste, readily soluble in Water, commercial samples frequently contain an undesirable excess of alkali in the form of Sodium Carbonate. **Zinc Lactate**, a white crystalline powder, or in glistening needle-shaped crystals, it has been used internally, in doses of 1 to 3 grains = 0.06 to 0.2 gramme, four or five times a day in the treatment of epilepsy.

CALCI LACTAS—White, mammillated tufts, or as a white odourless powder. Soluble 1 in 10 of Water, but solubility varies with the age of the preparation. Insoluble in Ether. Given in rachitis and scrofula, and in chilblains. As it increases the coagulability of the blood, it is given in hæmophilia, and before surgical operations on those who bleed unduly.

Dose.—3 to 10 grains = 0.2 to 0.65 gramme. Also combined with Ferri Lactas.

Foreign Pharmacopœias—Official in Belg., Ital. and Span.

FERRI LACTAS.—Pale greenish white crusts consisting of some needle-shaped crystals, or as a crystalline powder. Odourless when quite pure, but usually possessing a mild, peculiar odour and sweet, ferruginous taste. It should be kept in well stoppered bottles, as it tends to oxidise on exposure to air.

Solubility.—1 in 40 of cold Water, 1 in 12 of hot Water.

Dose.—5 to 15 grains = 0.32 to 1 gramme. In the form of a cachet or as a syrup.

Foreign Pharmacopœias—Official in all except U.S.

Tests.—Aqueous solutions of the salt give a deep blue precipitate with Potassium Ferricyanide, and a light blue precipitate with Potassium Ferrocyanide Solutions. When dissolved in diluted Sulphuric Acid and gently warmed after the addition of a little Potassium Permanganate the odour of Aldehyde is evolved.

1 gramme of the salt moistened with Nitric Acid and carefully ignited should leave a residue of Ferric Oxide amounting to not less than 0.27 gramme, indicating 27.0 per cent of Iron Oxide. This residue should not be alkaline in reaction to Litmus paper.

25 c.c. of a 1 in 50 aqueous solution of the salt after being boiled for a few minutes with 5 c.c. of diluted Sulphuric Acid, the Iron precipitated with an excess of Potassium Hydroxide Solution and filtered, the filtrate when boiled with a few drops of Potassic cupric Tartrate Solution should afford no reddish precipitate, indicating the absence of Sugar.

On trituration of a portion of the salt with strong Sulphuric Acid, no disagreeable odour should be evolved, no gas should be disengaged, nor should the mixture assume a dark colour after standing for some time. These tests show respectively the absence of fatty acids, Carbonates, and readily carbonisable organic impurities. A 1 in 50 aqueous solution of the salt should not produce more than a whitish opalescence with either Lead Acetate Solution or with Hydrogen Sulphide Solution after acidification with Hydrochloric Acid, indicating the absence of more than traces of Chloride, Citrate, Malate, Sulphate, Tartrate, and of heavy metals.

A 1 in 50 aqueous solution acidulated with dilute Nitric Acid shall yield no reaction with either Silver Nitrate Solution or with Barium Nitrate Solution, affording additional evidence of the absence of Chlorides and Sulphates.

The salt on being strongly heated evolves a caramel like odour, when more strongly heated gives off dense white fumes and finally leaves a brownish-red residue.

ACIDUM NITRICUM.

NITRIC ACID.

FR., ACIDE AZOTIQUE, GER., SALPETERSÄURE, ITAL., ACIDO NITRICO
CONCENTRATO, SPAN., ACIDO NITRICO

A clear, colourless fuming liquid, which evolves characteristic

choking fumes, and possessing even in diluted solutions a strongly acid and corrosive action.

It may be prepared by the decomposition of a Nitrate, generally Sodium or Potassium Nitrate by Sulphuric Acid.

It is officially required to contain 70 p.c. by weight of Hydrogen Nitrate, HNO_3 , eq. 62.58.

It should be preserved in well stoppered bottles, and in a cool place.

Medicinal Properties. It is strongly corrosive, and is applied as a caustic to warts, phagedenic sores, chancre, and condylomata, by means of a pointed glass rod. When diluted it is refrigerant, a stomachic tonic and cholagogue, and if very much diluted forms a drink in febrile diseases, and is used also as an injection to dissolve phosphatic calculi when of small size.

An acid of greater strength, 'Fuming Nitric Acid' (sp. gr. 1.5), is sometimes employed as a caustic.

Incompatibles. Alcohol, Alkali, Carbonates and Sulphides, Ferrous Sulphate, Lead Acetate.

Official Preparations.—*Acidum Nitricum Dilutum* and *Acidum Nitro-hydrochloricum Dilutum*. Used in the preparation of *Acidum Phosphoricum Concentratum*, *Argentum Nitratum*, *Liquor Ferri Perchloridi Fortis*, *Liquor Ferri Pernitratum*, *Liquor Ferri Persulphatis*, *Liquor Hydrargyri Nitratis Acidus*, *Spiritus Ætheris Nitrosi*, *Unguentum Hydrargyri Nitratis*.

Antidotes.—In case of poisoning by Nitric Acid, the antidotes are Chalk, Magnesia, or Carbonated Alkalis, with White of Egg, Carron Oil, or Soap-suds; followed by enemata of Beef Tea and Brandy, with Tincture of Opium to prevent collapse; emollient drinks.

Foreign Pharmacopœias.—Official in Austr., sp. gr. 1.300, Dan. and Norw., sp. gr. 1.180, Dutch, sp. gr. 1.316, B., sp. gr. 1.300, Fr., sp. gr. 1.304; Ger., Jap. and Swed., sp. gr. 1.153; Hung., sp. gr. 1.310, Ital., sp. gr. 1.400, Mex., sp. gr. 1.42, Port., sp. gr. 1.300 to 1.330, Russ., sp. gr. 1.200, Span., sp. gr. 1.31, Swiss, sp. gr. 1.151, also *Acidum Nitricum Fumans*, sp. gr. 1.45 to 1.5, U.S., sp. gr. 1.403 at 25° C (77° F.). Austr., Dan., Ger., Jap. and Norw., also *Acidum Nitrico-nitrosum*, sp. gr. 1.48 to 1.50.

Fr., Ger., Jap., Russ., Swed. and Swiss have an *Acidum Nitricum Crudum*.

Tests. Nitric Acid has a specific gravity of 1.420; the *B.P.* states 1.42, the *U.S.P.* about 1.403 at 25° C (77° F.); the *P.G.* 1.153. The boiling point should be about 120° C (250° F.); the *B.P.* states 121° C (250° F.); the *U.S.P.* 120.5° C (248° F.). It evolves dense red fumes when brought into contact with metallic Copper, it produces a dark brown coloration at the junction of the two liquids when a solution of the Acid is poured carefully upon a cooled mixture of Ferrous Sulphate and Sulphuric Acid, and when neutralised discharges the colour of a Sulphuric Acid Solution of Indigo, rapidly when warmed. The Acid is officially required to contain 69.46 p.c. of absolute Nitric Acid as indicated by titration with Volumetric Sodium Hydroxide Solution. As in the case of Lactic Acid, so here it would appear as if the atomic weights official in *B.P.* 1885 were employed in determining the number of c.c. of Volumetric Solution equivalent to the 1 gramme of Acid used, the yield of absolute Acid corresponding to 11.1 c.c. when so calculated being 69.93 p.c.

Calculating with the atomic weights of the 1898 Pharmacopœia, 11.2 cc would have been a nearer equivalent, which shows 70.08 pc of absolute Acid. The *USP* requires it to contain 68 pc by weight of absolute Acid, the *PG* 25 pc by weight.

The more generally occurring impurities are Arsenic, Copper, Lead and Iron, Bromic Acid or Bromine, Chlorides, Iodic Acid or Iodine, Sulphates and mineral residue. These are covered in the *BP* by the customary elastic expression, it should yield no characteristic reaction with the tests mentioned in the Appendix for those substances. Both *USP* and *PG* indicate the dilutions of the Acid to be employed, and in particular cases also the quantity of reagent to be used. The *USP* gives the modified Gutzeit test for Arsenic, and the time-limit test for heavy metals. A standard of 5 parts per million by weight is suggested (*CD* '08, 1795) for Arsenic, but 20 parts per million (by weight) is suggested for Lead, on account of the property of dissolving Lead from glass which this acid possesses. *PG* gives a special test for Iron, with Potassium Ferrocyanide Solution. In testing for Iodic Acid it will be noticed that the *USP* uses metallic Tin for the reduction, and the *PG* metallic Zinc.

The Pharmacopœias differ in the amount of mineral residue permitted. The *BP* states first that it shall yield no residue, and then 'not more than 0.005 pc', the *PG* omits all reference to the residue remaining after evaporation, whilst the *USP* states explicitly that 10 cc on evaporation to dryness, and further heated to 110° C (230° F), should leave no appreciable residue.

Modified Gutzeit Test—5 cc of a 1 in 10 aqueous solution of Nitric Acid should not respond to the modified Gutzeit's test for Arsenic, *USP*.

Potassium Ferrocyanide Solution—10 cc of a 1 in 10 aqueous solution should not be immediately turned blue on the addition of 0.5 cc. Potassium Ferrocyanide Solution, *PG*.

Chloroform.—Nitric Acid diluted 1-8, with a small piece of Zinc introduced for a short time, should not, on agitation with a small quantity of Chloroform, colour the Chloroform violet, *PG*.

The *USP* directs such a diluted Acid to be shaken with a few drops of Chloroform, which should remain colourless (absence of Iodine and Bromine) even after the introduction of a small piece of Tin (absence of Iodic and Bromic Acids).

Hydrogen Sulphide Solution—An aqueous 1 in 6 solution nearly neutralised with Ammonia Solution shall not be altered by the addition of Hydrogen Sulphide Solution, *PG*. When neutralised with Ammonia Solution and diluted with Distilled Water 1 to 20, the Acid should not respond to the time limit test for heavy metals.

Barium Chloride or Nitrate Solution—A 1 in 6 aqueous dilution of the Acid should not become more than opalescent within 5 minutes with Barium Nitrate Solution, *PG*. A separate portion of a 1 in 10 aqueous dilution should be unaffected by Barium Chloride Test Solution, *USP*.

Silver Nitrate Solution—A 1 in 6 aqueous dilution should not be affected by Silver Nitrate Solution, *PG*. A 1 in 10 aqueous dilution should be unaffected by Silver Nitrate Test Solution, *USP*.

Volumetric Determination—One gramme neutralises 11.1 cc of the Volumetric Sodium Hydroxide Solution, *BP*, 5 cc should require 22.9 cc of the Normal Volumetric Potassium Hydroxide Solution, *PG*. The *USP* directs that 3 cc of Nitric Acid be accurately weighed, diluted to 50 cc with Water and titrated with Normal Volumetric Potassium Hydroxide Solution,

using Methyl Orange Solution as an indicator. The number obtained by multiplying the number of c.c. of Alkali used by 6.257 and dividing this product by the weight of Acid taken represents the percentage of absolute Acid present.

Preparations.

ACIDUM NITRICUM DILUTUM. Diluted Nitric Acid

A clear, colourless acid liquid, prepared by diluting 3 fl. oz. and 7 fl. drms. of Nitric Acid with Distilled Water to make 20 fl. oz.

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

5 minims contain about 1 minim of strong Acid

Prescribing Notes.—*Usually diluted with Water or with bitter infusions and Tincture of Orange.*

Foreign Pharmacopœias.—Official in Belg., sp. gr. 1.072; Dutch, sp. gr. 1.133; Hung., sp. gr. 1.067; Ital., sp. gr. 1.1; Russ., sp. gr. 1.096; Fr., Jap. and Swiss, sp. gr. 1.056; U.S., sp. gr. 1.054 at 25° C (77° F). Not in the others, Dan., Norw. and Swed., see Acidum Nitricum.

Tests.—Diluted Nitric Acid has a specific gravity of 1.105; the U.S.P. states 1.054 at 25° C (77° F); the P.G. does not make a diluted Acid. It is officially required to contain 16.89 p.c. w/w of Hydrogen Nitrate as indicated by titration with Volumetric Sodium Hydroxide Solution; 1 gramme neutralising 2.7 c.c. The number of c.c. required to neutralise 1 gramme of the diluted Acid seems to have been based on the atomic weights official in the B.P. 1885, and using these latter would calculate to 17.01 p.c. of Hydrogen Nitrate. Each gramme should require for neutralisation 2.8 c.c. of the Volumetric Solution would have been more consistent with the statement of strength (17.1 p.c.) given in the official description of the diluted Acid.

As Nitric Acid (B.P.) is employed in its preparation, the diluted Acid is naturally required to answer the principal characteristic qualitative tests for Nitric Acid, and also be free from the impurities mentioned under the concentrated Acid.

ACIDUM NITRO-HYDROCHLORICUM DILUTUM. Diluted Nitro-Hydrochloric Acid

Nitric Acid, 3; Hydrochloric Acid, 4, Distilled Water, 25.

A clear, colourless, or pale yellow liquid, possessing a strong acid taste and faint chlorinous odour.

B.P. directs the Acids to be mixed with the Water and kept for 14 days before use; but scarcely any action takes place between the diluted Acids, free Chlorine and Nitrous Acid existing only in traces.

When the strong Acids were mixed, and after 8 days diluted, the resulting fluid liberated about fifty times as much Iodine from Potassium Iodide Solution as the B.P. preparation.

Medicinal Properties.—Cholagogue and gastric tonic. Externally as a lotion or bath, as well as by stomach administration for tropical enlargement and chronic congestion of the liver. Internally also in biliousness, in oxaluria, and in torpid conditions of stomach, intestinal glands and liver; and in catarrhal jaundice.

Nitro-Hydrochloric Acid Bath—Mix 8 oz by measure of Diluted Nitro-Hydrochloric Acid with 1 gallon of Water, temperature 96° or 98° F. Let a flannel roller of ten or twelve inches wide, and sufficient to encircle the body twice, be soaked in the fluid and then wrung, so as to remain only damp. Apply this instantly to the body, covering it with a piece of oiled silk to avoid damping the dress. It should be worn constantly, but should be changed, soaked, and wrung morning and evening. Glass, glazed earthenware, or wooden vessels should be used. Sponges and towels to be kept in Water to prevent them corroding.

The St Thomas's Hospital employ $\frac{1}{2}$ oz of the Diluted Nitro Hydrochloric Acid to the gallon for a full size bath of 25 to 30 gallons, and this has been incorporated in the *B.P.C.*

Dose—5 to 20 minims = 0.3 to 1.2 c.c.

Prescribing Notes—Usually diluted with Water and given with Tincture of Gentian or Tincture of Orange, and Tincture of Nux Vomica.

16 minims equal $1\frac{1}{2}$ minims of Nitric Acid and 2 minims of Hydrochloric Acid.

Incompatibles—Alkalis, Carbonates, Sulphides, salts of Silver and Lead.

Antidotes—See Acidum Nitricum.

Foreign Pharmacopœias—U.S. orders the undiluted—Nitric Acid, 18, Hydrochloric Acid, 82, also the diluted—Nitric Acid 4, Hydrochloric Acid, 18, Water, 78.

Norw., Nitric Acid, 1, Hydrochloric Acid, 2. By weight. *Syn* Aqua Regia.

Dublin Pharmacopœia was—Nitric Acid, 1, Muriatic Acid, 2.

Not in the other Pharmacopœias.

Tests—Diluted Nitro-Hydrochloric Acid has a specific gravity of about 1.07. 1 gramme should require for neutralisation about 2.5 c.c. of Volumetric Sodium Hydroxide Solution. When mixed with Potassium Iodide Solution, Iodine is liberated, which is readily recognised by the colour produced with Starch Solution.

ACIDUM OLEICUM.

OLEIC ACID

FR., ACIDE OLLIQUE, GER., OLEINSAURE, ITAL. AND SPAN., ACIDO OLEICO.

A pale brownish-yellow, oily liquid, which has a tendency to become rancid and to darken in colour on exposure to light and air. Pure Oleic Acid is represented by the formula $\text{HC}_{18}\text{H}_{33}\text{O}_2$, eq. 280.14, but the commercial article is usually not quite pure.

It is prepared from the Olein of fats, in which it exists as a Glyceryl ester, by saponification or hydrolysis, the former being accomplished by the Hydroxides of the fixed alkalis, the latter by the influence of superheated steam. After saponification the Oleate formed is decomposed by a mineral Acid.

It should be preserved in well-stoppered, dark amber-tinted glass bottles.

Solubility.—Mixes in all proportions with Alcohol, Chloroform, Ether, Benzol, Oil of Turpentine, and fixed Oils. Insoluble in Water.

Medicinal Properties.—Used in pharmacy for dissolving various metallic oxides and the alkaloids Morphine, Aconitine,

Atropine, Cocaine, and Veratrine, the oleates thus formed are more readily absorbed than ointments made with fats, oils, or paraffins

Official Preparation—Hydrargyri Oleas. Used in the preparation of Unguentum Atropine, Unguentum Atropine, Unguentum Cocaine, and Unguentum Veratrine. Of Mercuric Oleate, Unguentum Hydrargyri Oleatis.

Foreign Pharmacopœias—In Jap. (Acidum Oleicicum), sp. gr. about 0.9, M.P. 4.5 to 5°C. (40 to 41°F.), U.S. sp. gr. 0.895 at 25°C. (77°F.) Not in the others.

Tests.—Oleic Acid has a specific gravity of from 0.890 to 0.910; the *U.S.P.* states 0.895 at 25°C. (77°F.). The solidifying point is 4.5 to 5°C. (40 to 41°F.), subsequently melting at 13.3 to 15.5°C. (56 to 60°F.), the *U.S.P.* states that it becomes semi-solid when cooled to from 0 to 4°C. (32 to 39.2°F.) and congeals to a whitish solid on further cooling. It has a peculiar characteristic odour. The acid may be determined by titration with Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 gramme of the acid should require about 3.5 cc. of Volumetric Sodium Hydroxide Solution, indicating 98.0 p.c. of absolute Oleic Acid. No volumetric assay is included in either the *U.S.P.* or *B.P.*

The most likely impurities are fixed oils, Stearic and Palmitic Acids. The solubility in Alcohol (90 p.c.) detects the presence of fixed oils; though the solubility is not specifically mentioned as a test for these in the *B.P.*; advantage is taken of the comparative insolubility of Lead Stearate and Lead Palmitate in Ether to use this agent as a test for the presence of these two Acids.

The *U.S.P.* states that heated to a temperature of 95°C. (203°F.) decomposition commences, on heating to a higher temperature it is entirely dissipated.

The Acid is not official in the *P.G.*

Solubility in Alcohol. The *U.S.P.* states that an alcoholic solution of the Acid should have a feebly acid reaction on Litmus paper, and that equal volumes of Acid and Alcohol should form a clear solution at 25°C. (77°F.) without the separation of oily drops, indicating absence of fixed oils.

Lead Acetate Solution.—A weighed quantity of 1 gramme of Oleic Acid is dissolved in about 20 cc. of Alcohol (94.9 p.c.) and 2 drops of Phenolphthalein Solution added. The mixture is warmed and saponified by the addition drop by drop of a Solution of Sodium Hydroxide (1-4) until the Acid is neutralised, and the mixture after agitation remains a permanent red colour. Then gradually add Acetic Acid until the red tint is discharged, filter, and mix 10 cc. of the filtrate with 10 cc. purified Ether. This solution should not become any more than slightly turbid with 1 cc. of Lead Acetate Solution, indicating the absence of notable quantities of Stearic and Palmitic Acids, *U.S.P.*

An almost identical test is employed by the *B.P.* for the detection of more than traces of these acids.

Not Official.

ACIDUM STEARICUM Stearic Acid $\text{HC}_{18}\text{H}_{35}\text{O}_2$, eq. 282.14.—A hard white glistening solid, possessing very little odour or taste. It is an organic acid appearing commercially in a more or less impure form as Stearine. It is a monobasic acid. It is insoluble in Water, soluble in Alcohol (90 p.c.), and readily dissolves in Ether.

Tests.—Stearic Acid melts at 69.2°C. (156.6°F.). This melting point is given in the *U.S.P.*, which further states that the commercial acid should have a

melting point not lower than 56°C (132°F). The solidifying point of the commercial acid should not be lower than 54°C (129°F). It may be determined volumetrically by titration of a weighed quantity with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality 1 cc of the Volumetric Solution corresponding to 0.28214 gramme of pure Stearic Acid. Hydrocarbon oils and unsaponified fat are the more generally occurring impurities. If present in more than traces both are readily detected by the saponification test. If 1 gramme of the acid be saponified by boiling with a solution of 0.5 of a gramme of exsiccated Sodium Carbonate in 30 cc of Water, the resulting liquid should not be more than opalescent. It should leave no weighable residue when ignited with free access of air, indicating the absence of mineral matter, Soap, etc.

Not Official

ACIDUM OSMICUM

OSMIC ACID



Fr, ACIDE OSMIQUE, Ger, Osmiumsäure

A pale yellow crystalline substance giving off an excessively irritating vapour, which attacks the eyes and nostrils. It is more convenient in the form of 1 p.c. aqueous solution, which must be carefully protected from dust or organic matter which will reduce it and form a black deposit.

Solubility—1 in 17 Water. Should not be dissolved in Alcohol or Ether, as decomposition ensues.

Medicinal Properties—4 to 10 minims of a 1 p.c. aqueous solution of Osmic Acid or Potassium Osmate have been injected hypodermically for sciatica and other forms of neuralgia.

5 to 10 minims in two or three separate injections in certain forms of neuralgia.—L '99, ii 1250, '03 ii 970

In trigeminal neuralgia the main branches of the fifth nerve were exposed, and a few drops of a 2 p.c. solution injected at several points.—M P '04, ii 470. Injected into Gasserian ganglion.—L '07, ii 1603

Used as 1 p.c. aqueous solution for fixing and staining in histological work. Fat and nerve tissues are blackened by it.

Foreign Pharmacopœias—Official in Mex

Not Official

ACIDUM OXALICUM



Fr, ACIDE OXALIQUE, Ger, Oxalsäure, Ital, ACIDO OSSALICO

This is noticed here rather as a poison than a medicine, although it has been used in America in the treatment of amenorrhœa, and as a sedative in acute cystitis (*T G* '91, 164) in $\frac{1}{2}$ grain doses every four hours. It is used in households for cleaning brass, and removing ink stains, iron moulds, etc. It has been mistaken for Epsom Salts, which it somewhat resembles. Murrell states that death has occurred from 2 drms, but recovery from $\frac{1}{2}$ oz.

Antidotes—Chalk, Lime, or Whitening are given freely in Water. Saccharated Solution of Lime may be given in drim doses, frequently repeated, also emollient and stimulant drinks.

Foreign Pharmacopœias—Official in Mex and Port

ACIDUM PHOSPHORICUM CONCENTRATUM.

CONCENTRATED PHOSPHORIC ACID

FR., ACIDE PHOSPHORIQUE OFFICINAL., GER., PHOSPHORSÄURE, LIQ.,
ACIDO FOSFORICO SPAN., ACIDO FOSFORICO

A clear, colourless, and odourless syrupy liquid, containing 66·3 p.c. of Hydrogen Orthophosphate, H_2PO_4 , eq. 97·32

It is prepared from Phosphorus by oxidation. The official method is stated to be the oxidation, by means of Nitric Acid, of the aqueous solution of the residue from the atmospheric oxidation of Phosphorus. The more general method of production is the direct oxidation of Phosphorus by means of Nitric Acid.

It should be preserved in well stoppered glass bottles, preferably of an amber tint.

Medicinal Properties. This concentrated Acid is used in making phosphatic preparations. Only given internally in the diluted form. See Acidum Phosphoricum Dilutum.

Official Preparations. Acidum Phosphoricum Dilutum. Used in the preparation of Acidum Hydrobromicum Dilutum, Ammonii Phosphas, Syrupus Calcii Lactophosphatis, Syrupus Ferri Phosphatis, and Syrupus Ferri Phosphatis cum Quinina et Strichnina.

Foreign Pharmacopœias. Official in Austr., sp. gr. 1·12 (20 p.c.), Belg., 1·056 to 1·057 (10 p.c.), Fr. and Ital., sp. gr. 1·319 (50 p.c.), Dutch, sp. gr. 1·153 (25 p.c.), Ger. and Russ., sp. gr. 1·151 (25 p.c.), Jap., Hong., sp. gr. 1·120 (20 p.c.), Mex., 1·34, Port., sp. gr. 1·880. Spain, sp. gr. 1·35 (50 p.c.), U.S., sp. gr. not below 1·707 at 25° C. (77° F.) (85 p.c.). Not in the others.

Tests.—Phosphoric Acid has a specific gravity of 1·5; the U.S.P. acid 1·707 at 25° C. (77° F.), the P.G., 1·154. A white crystalline precipitate is produced when its diluted neutralised aqueous solution is mixed with Magnesium Ammonio sulphate Solution. A yellow precipitate soluble in Ammonia Solution is produced when its diluted aqueous solution acidified with Nitric Acid is treated with Ammonium Molybdate Solution containing free Nitric Acid. A canary yellow precipitate readily soluble in Ammonia Solution, and in cold dilute Nitric Acid is produced when Silver Ammonio-nitrate Solution is added to a dilute neutralised aqueous solution of this Acid. The Acid is officially required to indicate when assayed according to the Loid process given below, 66·3 p.c. of Hydrogen Orthophosphate; the U.S.P. Acid should contain 85 p.c. by weight of absolute Orthophosphoric Acid, the P.G. contains 25 p.c. of pure Acid. The process of determination adopted by the B.P. is a gravimetric one, and consists in converting the Phosphoric Acid into a Lead salt by means of Lead Oxide, evaporating and heating the residue to dull redness. A weighed quantity of 1 gramme of the Acid treated with 2·5 grammes of finely powdered Lead Oxide is required to yield a residue weighing 2·98 grammes. As pointed out in the 17th Edition of the Companion the percentage acidity of Phosphoric Acid may be conveniently determined by titration with standard alkali, using Phenolphthalein Solution as an indicator; the change of colour takes

place when two-thirds of the Hydrogen is replaced by alkali metal, with Methyl Orange Solution as the indicator, neutrality is reached with half the quantity of alkali, with Litmus Solution the end reaction is too indefinite. The process adopted by the *USP* is volumetric, Normal Volumetric Sodium Hydroxide Solution being employed and Phenolphthalein Test-solution used as an indicator of neutrality. A definite weight of the Acid is diluted with Water, and the quantity removed for the determination is then diluted with about an equal quantity of a cold aqueous solution containing 5 grammes of Sodium Chloride. The *PG* gives neither a gravimetric nor a volumetric process of determination.

The more generally occurring impurities are Meta- and Pyrophosphoric Acids, Phosphorous Acid, Arsenic, heavy metals (*e.g.*, Copper, Iron, and Lead), Calcium, Chlorides, Sulphates, Phosphates, and Nitrates. The *BP* are content to include most of these under the general expression it 'shall yield no characteristic reaction,' etc.

The *BP* and the *USP* employ Albumen Solution for proving the absence of Meta-phosphoric Acid, and Tincture of Ferric Chloride for proving the absence of Pyro phosphoric and Meta phosphoric Acids. The tests in each Pharmacopœia are virtually the same, requiring that when diluted with 5 volumes of Water no precipitate should be produced on the addition of Albumen Solution, nor should any precipitate be formed, even after several hours, on the addition of an equal volume of Tincture of Ferric Chloride.

Individual tests for Arsenic are given by the *USP* and *PG*, the former employing the modified Gutzeit's test, the latter Solution of Stannous Chloride. Both these Pharmacopœias employ Hydrogen Sulphide for the detection of heavy metals, the *USP* adopting the lime limit test. In testing for Nitrates both *USP* and *PG* employ equal volumes of Phosphoric and Sulphuric Acids, the former using a crystal of Ferrous Sulphate, the latter a Solution of Ferrous Sulphate. In testing for Phosphates *USP* employs a mixture of three parts of Alcohol (94.9 p.c.) and one of Ether, whilst *PG* uses Alcohol (90 p.c.) alone. No such test appears in the *BP*. A test for the absence of Silica is included in the *BP*, but not in *USP* or *PG*.

The *BP* employs Mercuric Chloride Test solution as a reagent for the detection of Phosphorous Acid, the *USP* both Mercuric Chloride Test-solution and Silver Nitrate Solution, the *PG* Solution of Silver Nitrate and gentle warming. When made alkaline with Ammonia Solution it should not give (even after long standing) a crystalline precipitate of Ammonio-magnesium Phosphate, indicating absence of Magnesium, which is present to a considerable extent in some commercial samples.

The acid when diluted with Water should yield no precipitate when allowed to remain at rest for some time, indicating the absence of Silica.

Mercuric Chloride.—The *BP* requires that, if the Acid be diluted, mixed with an equal volume of Mercuric Chloride Solution and heated, no precipitate should form, the *USP* gives 1 c.c. of Acid diluted with 5 c.c. of Water, but no

specified quantity of reagent, a similar dilution when gently warmed and with the addition of a few drops of Silver Nitrate T.S. should not be blackened, indicating the absence of Phosphorous Acid.

Silver Nitrate Solution. Phosphoric Acid (diluted with 5 volumes of Water, *U.S.P.*) should yield no precipitate with Silver Nitrate T.S., indicating the absence of Chlorides, *P.G.* and *U.S.P.* (not even on warming, indicating the absence of Phosphorous Acid, *P.G.*)

Barium Chloride or Nitrate Solution. A diluted Acid (1:1, *P.G.*) should be unaffected by Barium Nitrate Solution, *P.G.*, indicating the absence of Sulphates.

0.1 cc. of the Acid diluted with Water to 7 cc. should not afford a cloudy mass or precipitate within 30 seconds on the addition of 1 cc. of Barium Chloride Test solution, *U.S.P.*

Ammonium Oxalate Solution. A 1:1 aqueous dilution of the Acid, after the addition of excess of Ammonia Solution should not be changed by the addition of Ammonium Oxalate Solution, indicating the absence of Calcium, *P.G.*

Hydrogen Sulphide. The Acid should not be affected by Hydrogen Sulphide Solution, *P.G.*, and 10 cc. of a 1:20 dilution should not respond to the time limit test for heavy metal.

Modified Gutzzeit's Test. 5 cc. of a 1:10 aqueous dilution should not respond to the modified Gutzzeit's test for Arsenic, *U.S.P.*

Stannous Chloride Solution. The *P.G.* requires that a mixture, 1 cc. of Phosphoric Acid with 3 cc. Stannous Chloride Solution, should not assume a dark colour in the course of an hour, *P.G.*

Alcohol or Alcohol and Ether. In a mixture of 1 cc. Phosphoric Acid, 3 cc. Alcohol (94.9 pc) and 1 cc. Ether there should be no turbidity (absence of Phosphates) *U.S.P.* The *P.G.* directs that a mixture of the Acid with 4 volumes of Alcohol (90 pc) should remain clear.

Ferrous Sulphate. The *U.S.P.* requires that no white precipitate or colour should appear around a crystal of Ferrous Sulphate dropped into a cooled mixture of 1 cc. Phosphoric Acid and 1 cc. Sulphuric Acid. No coloured zone should be formed when 1 cc. Ferrous Sulphate solution is poured as a layer on a mixture of 2 cc. Phosphoric Acid and 1 cc. Sulphuric Acid, *P.G.*

Volumetric Determination. If 10 grammes of Phosphoric Acid be diluted with Water to measure 100 cc., then 9.73 cc. of this, diluted with 10 cc. of cold saturated aqueous solution containing 5 grammes of Sodium Chloride, should require 17 cc. of Normal Volumetric Potassium Hydroxide Solution for neutralisation (each cc. corresponding to 5 pc. of absolute Phosphoric Acid), Phenolphthalein T.S. being used as indicator, *U.S.P.*

Preparation.

ACIDUM PHOSPHORICUM DILUTUM. DILUTED PHOSPHORIC ACID.

A clear colourless liquid possessing an acid taste and strong acid reaction towards blue Litmus paper. It is prepared by diluting 3 of Concentrated Phosphoric Acid with Distilled Water to make 20.

Medicinal Properties.—Tonic and refrigerant, hamatinic and anhydrotic; diuretic in the phosphatic diathesis. Given with Calcium Phosphate in rickets. Quenches the craving for fluids in diabetes.

Used as a partial substitute for organic acids in cooling drinks and acidulated waters.

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

Prescribing Notes.—Usually largely diluted with Water, and given with some other and aromatic mixtures and syrups; should not be mixed with the Syrup of Iron Pyrophosphate, as the mixture becomes solid.

Incompatibles.—Lime Water and all alkalis

Foreign Pharmacopœias—Official in Norw and Port, 13·8 p.c., Russ., 12·5 p.c., Fr., Jap., Mex., Swed., Swiss and U.S., 10 p.c.

Tests—It has a specific gravity of 1·08, the *USP* diluted acid has a specific gravity of 1·057, the *PG* does not include a diluted acid. It is officially required to contain 13·8 p.c. w/w of Hydrogen Orthophosphate as gravimetrically determined by converting 1 gramme into a Lead salt by the addition of 0·5 of a gramme of Lead Oxide, heating to a dull redness, cooling and weighing the residue, which should amount to 0·6 gramme, the *USP* diluted acid is required to contain 10 p.c. w/w of absolute Ortho-phosphoric Acid as determined volumetrically by titration with Normal Volumetric Potassium Hydroxide Solution, using Phenolphthalein Test-solution as an indicator of neutrality. The quantity (4·87 gramme) of the diluted Phosphoric Acid used for the titration is diluted with 5 c.c. of a cold saturated Sodium Chloride Solution. Concentrated Phosphoric Acid answering the official description is used in the preparation of the diluted acid, and it is therefore required to answer the tests and to be free from the impurities given under the Concentrated Acid.

Not Official

ACIDUM PICRICUM

$C_6H(NO)_3OH$, eq 227·44

PICRIC ACID CARBAZOIC ACID TRINITROPHENOL.

Pale yellow crystalline scales, prepared by the action of hot Nitric Acid on Phenol sulphonic Acid.

Picric Acid, Picrates, and Mixtures of Picric Acid when in process of manufacture or when kept, conveyed, imported, or sold for any purpose, come within the Explosives Act 1875, except when it is mixed with not less than half its own weight of Water. For the dealer in these substances there are special conditions as to storing, etc.

With Ammonia, Potassium Hydroxide and Sodium Hydroxide it forms crystalline salts, which are explosive.

Solubility—1 in 90 of Water, 1 in 10 of Alcohol (90 p.c.)

Medicinal Properties—A solution (1 or 2 p.c.) of Picric Acid has been recommended as an application to scalds and burns, and also in acute eczema.—*B M J* '96, ii 651 and 1826, '97, i 331 and 457, '99, i 1152, *L* '03, ii 640, 799.

It has also been given in $\frac{1}{2}$ to 1 grain doses as a bitter tonic.

Specially useful as a first dressing in burns, has all the advantages of boric acid, plus that of relieving pain.—*B M J* '07, ii 524.

Solution for Removal of Picric Acid Stains. Sodium Benzoate, 1; Boric Acid, 1, Water, 100.

A saturated aqueous solution is a delicate test for the presence of Albumen in fluids, even in very dilute solutions a white cloud is formed at the junction of the two liquids, and in stronger solutions the Albumen is precipitated. Used in histological work.

Foreign Pharmacopœias—Official in Fr., Jap. and Mex. Not in the others.

Tests—The pure Acid melts at 122° C (241·6° F). At a higher temperature it partially sublimes, and boils, giving off bitter yellow suffocating vapours. It may be determined by titration with Volumetric Sodium Hydroxide Solution,

using Phenolphthalein Solution as an indicator of neutrality. 1 gramme of the Acid should require about 4.4 c.c. of the Volumetric Solution. A dark red liquid is produced when a solution of Picric Acid is boiled with a strong Solution of Potassium Cyanide. When boiled with a strong Solution of Calcium Hypochlorite it gives off picric acid, producing vapours. It should be free from mineral impurities, and from more than traces of Sulphates.

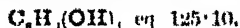
AMMONII PICRAS.—Yellow, odourless, glistening crystalline needles. Soluble 1 in 93 of Water 1 in 82 of Alcohol (90 p.c.) Given as a substitute for Quinine, also in exophthalmic goitre and malaria.

Dose.— $\frac{1}{2}$ to 1 gram = 0.016 to 0.06

Not Official

ACIDUM PYROGALLICUM.

PYROGALLIC ACID, PYROGALLOI.



FR., PYROGALLOL, GER., PYROGALLOL, ITAL., PYROGALLOLO.

Light, white crystalline tufts, which have a tendency to become coloured on exposure to strong light, more particularly in solution. The change is more rapid in alkaline solution. Usually prepared by heating Gallic Acid to 185° to 200° C. (365° to 392° F.)

It should be kept in well closed, dark amber-tinted glass bottles as far as possible from the light.

Solubility.—1 in 2 of Water, and measures 2 $\frac{1}{2}$, 9 in 10 of Alcohol (90 p.c.).

Medicinal Properties.—Escharotic, antiseptic, and disinfectant. Its use requires care.

Not more than 15 to 25 grains should be used in the twenty-four hours, as violent toxic symptoms may result from its absorption. *T. G.* '85, 59.

Used in the form of a 10 p.c. salve, and applied with a brush twice a day, it proved very useful in Hebra's wards in the treatment of psoriasis. The parts were then covered with cotton wadding or linen, and when very extensive were covered with flannel. *L. M. R.* '87, 377.

An ointment, Pyrogallin Acid 10, Starch 40, Vaseline 120, also a powder, Pyrogallin Acid 20, Starch 80, have been used for venereal ulcers. *L. M. R.* '82, 228, '84, 68.

Mixed with Colloodium Flexile, 40 grains to the oz., for psoriasis. — *T. G.* '88, 181.

Largely used in photography. It has also been used for bleaching the hair. 1 in 16 of Water is used with a Solution of Silver Nitrate (1 in 30 of Water).

To remove stains of Pyrogallin Acid, rub a little Ammonium Persulphate on the fingers and rinse with Water. *P. J.* '98, 1504a.

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Russ., Swed., Swiss and U.S. Not in Port. or Span.

Tests.—Pyrogallin Acid melts at 131° to 132° C. (267.8° to 269.6° F.), and sublimes at a higher temperature without leaving any mineral residue. Solutions of the acid gradually absorb Oxygen from the air becoming darker in colour, and this absorption is much accelerated in the presence of alkalis, the solutions then rapidly becoming dark brown or almost black. A freshly prepared dilute Solution of Ferrous Sulphate yields an indigo blue coloration with a solution of Pyrogallin Acid, and solutions of Ferric salts a brownish-red coloration; solutions of Mercuric, or Silver salts are rapidly reduced.

UNGUENTUM ACIDI PYROGALLICI (Jarsch's Ointment) —Pyrogallin Acid, 60 grains; Lard, 1 oz.

This has been incorporated in the B.P.C. as follows.—

Pyrogallin Acid, 12, Lard, 88

UNGUENTUM PYROGALLOL COMPOSITUM.—Pyrogallie Acid, 80 to 60 grains, Ichthyol, 80 grains, Salicylic Acid, 15 grains, Soft Paraffin, to 1 oz — *Middlesex*

Pyrogallol, 20 grains, Ammonium Sulpho-ichthyolate, 20 grains, Salicylic Acid in powder, 8 grains, Soft Paraffin to 1 oz — *London*

Unguentum Pyrogalloli Compositum (Unna)—Pyrogallie Acid, 5, Salicylic Acid, 2, Ammonium Ichthiosulphonate, 5, Yellow Vaseline, 88 — *Hager, and Pharm. Form*

This has been incorporated in the *B P C* under the title **Unguentum Acidi Pyrogallici Compositum** *syn* Unna's Compound Pyrogallol Ointment, employing Soft Paraffin

Unguentum Pyrogallol Oxydati—Oxidised Pyrogallie Acid, 5, Salicylic Acid, 5, Hydrous Wool Fat, *q s* to make 100 — *B P C*

UNNA'S PYROGALLIC PLASTER MULL—Contains 40 p c of the Acid, equal to $\frac{1}{2}$ grain in each square inch of surface

ACID PYROGALLIC OXIDISED (Pyraloxin)

Some attention has been directed to this drug by its recent employment in dermatological practice. It is a brownish black powder readily soluble in Water, prepared by oxidising Ammonium Pyrogallate in a current of air. It possesses no toxic properties, nor is it liable, as a rule, to excite any dermatitis. It has been employed (*Edin Med Jour* '05, 487) in the treatment of lupus erythematosus.

In the treatment of psoriasis it stands next to Chrysarobin in efficiency. It is unsuited for the acute or rapidly extending phase, but when the disease has come to a standstill, or is showing distinct evidences of retrogression, its advantages are said to be incontestable. It may be applied as an ointment ($\frac{1}{2}$ drm to the oz), made up with Vaseline, or with the addition of 10 grains of Salicylic Acid as a mordant. A cleaner method is to employ 10 parts dissolved in 20 of Benzol and 80 of Acetone.

In lupus erythematosus applied as an ointment. Zinc Oxide, 10, Kaolin, or terra silicia, 2, Oxidised Pyrogallie Acid, 5, Vaseline, 28.

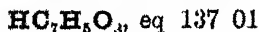
In infantile eczema of the face the most brilliant results have been obtained. It is applied to the reddened and irritable surface of the skin in the form of a thin coating of Lassar's Paste, to which 10 grains of Pyraloxin have been added.

For the eradication of ringworm of the scalp in children it is applied in the form of an ointment. Oxidised Pyrogallie Acid, 10 grains, Precipitated Sulphur, $\frac{1}{2}$ drm, Ammoniated Mercury, 15 grains, and Vaseline, 1 oz.

Eugallol (Pyrogallol mono-acetate), a brownish yellow thick syrupy liquid, **Gallacetophenone**, a yellowish brown powder, **Lenigallol** (Pyrogallol triacetate), a white powder, and **Saligallol** (Pyrogallol di salicylate), a resinous solid, are preparations which have received attention in the treatment of skin diseases.

ACIDUM SALICYLICUM.

SALICYLIC ACID



FR, ACIDE SALICYLIQUE, **GER**, SALICYLSÄURE, **ITAL**, ACIDO SALICILICO, **SPAN**, ACIDO SALICILICO

Colourless, odourless, prismatic crystals when prepared synthetically from Carboic Acid, but the Acid derived from the Oil of Wintergreen or of Sweet Buch, commonly called 'natural acid,' is usually supplied in large crystals possessing a yellowish or pinkish tint, and generally possesses a faint odour of Methyl Salicylate.

It may also occur as a light, white crystalline powder. It possesses at first a sweetish and subsequently an acrid taste.

Prepared by passing Carbonic Acid Gas into a mixture of Carboic Acid and Sodium Hydroxide at a high temperature, and decomposing the Sodium Salicylate with a mineral Acid, and subsequent purification; or by treating Oil of Wintergreen (*Gaultheria procumbens*), which is mainly composed of Methyl Salicylate, also Oil of Sweet Birch (*Betula lenta*) and *Andromeda leschenaultii* (a native of India), with a Solution of Potassium Hydroxide, and distilling it, decomposing the residue with Hydrochloric Acid, and purifying the Salicylic Acid by recrystallisation.

Salicylic may be sublimed, but there is almost certain to be some slight decomposition with liberation of Phenol.

Solubility. - About 1 in 550 of Water, 1 in 9 of boiling Water, 1 in $3\frac{1}{2}$ of Alcohol (90 p.c.), 1 in 11 of Alcohol (60 p.c.), 1 in 35 of Alcohol (15 p.c.), 1 in 2 of Ether; 1 in 55 of Chloroform; 1 in 120 of Olive Oil, 1 in 195 of Glycerin; 1 in 8 of Lard (at 180° F.). 20 grains Salicylic Acid are rendered soluble in a fl. oz. of Water by the addition of 25 grains of Borax or 10 grains of Potassium Citrate; but it is better to use Soda or Salicylate.

Medicinal Properties. Antiseptic and powerfully antipyretic; specific in acute rheumatism.

A good **preservative** of medicated solutions, such as Cocaine salts and Boric Acid, which are otherwise liable to fungoid growths, 1 in 1000 is sufficient for the purpose, but in the eye causes temporary smarting.

Used as a **lotion** (4 p.c.) in pruritus and urticaria, and some forms of eczema; as an **injection** (1 in 300) in the dysenteric diarrhoea of children; as an **ointment** (1 in 6) for pruritus (*Ringer*). With Zinc Oxide and Starch it is used as a '**dusting powder**' for infants.

In **collodion** form it is very useful when applied to hard and soft corns and warts. It softens and removes them.

The collodion is recommended in lupus. *P* 1 in '96.

The injection of Salicylic Acid in uterine cancer, recommended as a **palliative** method when the disease is too far advanced to admit of surgical extirpation. *P. J* (8) xxv 1219.

A solution in spirit, increased 1 p.c. daily from 1 to 6 p.c., applied to remove the stumps left after removal of papillomatous growths. It also forms a useful application dissolved in Sodium Sulpho ricinate solution. — *B. M. J.* '04, ii. 1221, 1224.

A daily application of a 10 p.c. solution of Salicylic Acid in Sulpho ricinate of Soda for pharyngo mycosis. This salt of soda seems to be the vehicle with which Salicylic Acid can be incorporated so as to be as unirritating as possible. — *B. M. J.* '07, i 1155, *L.* '07, i. 1248.

Recommendation of the Departmental Committee appointed to inquire into the use of preservatives and colouring matters in food. Salicylic Acid be not used in greater proportions than 1 grain per pint in liquid food, and 1 grain per lb. in solid food, its presence in all cases to be declared. *L.* '01, ii. 1643, *J. S. C. I.* '01, 1228, *P. J.* '01, ii. 620, *C. D.* '01, ii. 880, *Analyst*, '01, 882.

Salicylic Acid as a preservative for foods has been stated to be injurious, for three reasons: (1) That it is antiseptic and antifermentative, and therefore liable to interfere with the digestive processes, (2) after absorption is apt to injure the general health, (3) that it is an irritant and apt to injure the mucous membrane of stomach and intestinal canal, but the results of experiments by Macalister and Bradshaw are opposed to these conclusions, and they contend that there is no justification for them. — *L.* '02, ii. 1644; '03, i 717.

Prescribing Notes.—On account of its slight solubility in Water, it is usually given internally in the form of Sodium Salicylate, which is readily soluble, and is less irritating to the mucous membrane. It is also given in combination with Bismuth and Lithium.

Dose.—5 to 20 grains = 0.32 to 1.3 grammes

Incompatibles—Spirit of Nitrous Ether, Iron salts

Official Preparations—Sodu Salicylas, Unguentum Acidi Salicylic Used in the preparation of Injectio Cocainæ Hypodermica, Liquor Atropinæ Sulphatis, and Salol. See also Bismuthi Salicylas

Not Official.—Collenplastrum Salicylatum, Collodium Salicylicum, Collodium Lacto salicylicum, Collodium Callosum, Salicylic dressings, Glycerinum Acidi Salicylici, Lotio Acidi Salicylici Co, Parogen Salicylatum, Pulvis Salicylicus cum Talco, Pulvis Talci Salicylicus, Salicylic and Creosote Plaster Mulla, Salicylic Acid Suet, Unguentum Acidi Salicylici, Vasolimentum Salicylicum, Salacetol, Aspirin, Acetylsalicylic Acid, Agathin, Glycosal, and Salitannal

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Jap, Ital, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests.—Salicylic Acid melts at 156° to 157° C (312.8° to 314.6° F), the *USP* states it begins to melt at 156° C (312° F) and is completely melted at 157° C (314.6° F), the *PG* that it melts at about 157° C (314.6° F). According to the *USP* it is gradually dissipated at a higher temperature than 157° C (314.6° F). The distinctive test is the violet colour imparted to an aqueous solution by Ferric Chloride Test-solution, and the production of the peculiar odour of Methyl Salicylate when a little of the Acid is warmed with Methyl Alcohol and strong Sulphuric Acid. The latter test is included in *USP* but not in *BP* or *PG*. The Acid is not officially required in this instance to 'afford when neutralised the reactions characteristic of Salicylates.' According to the *BP* the acid volatilises at 200 C (392° F) 'without decomposition,' but unless the Acid is very carefully heated there is almost certain to be some slight decomposition, with liberation of Phenol, which fact is noted in *PG*. The Acid may be readily determined by titration with Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. Each gramme of the Acid should require about 7.3 c.c. of the Volumetric Solution, indicating 100.0 p.c. of absolute Salicylic Acid. Neither the *BP* nor *PG* includes a volumetric method of determination.

The more generally occurring impurities are unconverted Phenol, isomers, or homologues of Salicylic Acid, organic impurities, Hydrochloric Acid, Carbolates or Sulphocarbolates, and mineral residue. The test for Phenol, viz., its extraction from alkaline solution with Ether, is practically the same in the *BP*, *USP*, and *PG*, the latter Pharmacopœia giving the relative quantities to be used in performing the test. The method adopted by each Pharmacopœia is compared in the small type below, under the heading of Ether. Isomers or homologues of Salicylic Acid are detected by their influence on the melting point and by the evaporation test. The *BP* evaporates an aqueous solution, whilst *PG* and *USP* employ an alcoholic one. The test is more severe when water is employed with excess of the sample, for as pointed

out (*P.J.* (3) xxi 478), the impurities are more soluble in this fluid than the Acid, it therefore tends to magnify the proportion of impurity on solution, and to separate it on evaporation. A filtered saturated aqueous solution prepared by shaking the acid with a small quantity of Water, when evaporated to dryness is officially required to leave a white residue free from any buff coloured fringe, which according to the *B.P.* indicates the absence of Iron, organic impurities and colouring matter. Concentrated Sulphuric Acid serves to detect organic impurities, the *B.P.* requiring no colour to be developed in 15 minutes, but making no reference to the relative proportions of the two acids, the *U.S.P.* and *P.G.* giving the latter but no time limit, the *U.S.P.* employs about 0.5 of a gramme of Salicylic Acid in 10 c.c. of Sulphuric Acid, the *P.G.* requires one part of Salicylic Acid should be dissolved by 6 parts of Sulphuric Acid without coloration. A test for Chlorides is included in *U.S.P.* and *P.G.* The *U.S.P.* allows not more than 0.6 p.c. and the *P.G.* no weighable residue; the *B.P.* makes no mention of mineral residue.

The test distinguishing Salicylic Acid from Carbolates and Sulphocarbolates by means of Solution of Uranium Nitrate is peculiar to the *B.P.*; but this reaction would not detect the presence of either of the latter in a sample of Salicylic Acid. It depends upon the assumption that Uranium Nitrate Solution produces a yellowish-brown precipitate in solutions of the acid not weaker than 1 p.c., whereas solutions of Carbolates and Sulphocarbolates are not precipitated.

Evaporation. In the *P.G.* and *U.S.P.* an alcoholic solution of the Acid (1 10, *P.G.*, saturated, *U.S.P.*) is allowed to evaporate spontaneously in a place protected from dust, when a perfectly white residue should remain.

Ether. If 1 gramme of Salicylic Acid be dissolved in excess of cold Sodium Carbonate Solution and the liquid agitated with an equal volume of Ether, and the ethereal solution be allowed to evaporate spontaneously, the residue, if any, should be free from the odour of Phenol, *B.P.* and *U.S.P.* The *P.G.* test gives as quantities 0.5 gramme of Acid dissolved in 10 c.c. of a 1-10 Sodium Carbonate Solution.

Silver Nitrate. The *P.G.* and *U.S.P.* require that an alcoholic Solution of Salicylic Acid (1 10, *P.G.*, 1 20, *U.S.P.*) should be unaffected by Silver Nitrate T.S. after the addition of a few drops of Nitric Acid, indicating the absence of Chlorides.

Preparation.

UNGUENTUM ACIDI SALICYLICI. SALICYLIC ACID OINTMENT

Salicylic Acid, 1; White Paraffin Ointment, 49.

Foreign Pharmacopœias. Official in Mex. (Ponada de Acido Salicilico), Acid 1, Alcohol 2, Vaseline 9, Austr., Sodium Salicylatum (see below). Not in the others.

SODIUM SALICYLATE. *See* SODII SALICYLAS.

Not Official.

COLLEMPASTRUM SALICYLATUM (*Austr.*) - Collemplastum adhaesivum mass, 100; Acid Salicylic, 4, Petroleum Ether, 20.

COLLEMPASTRUM ADHÆSIVUM (*Austr.*) - Resin Oil, 6, purified alcohol India-rubber, 10; Petroleum Ether, 45, allow to stand with frequent

agitation until dissolved, add Balsam Copaiba and Colophony Resin, of each, 4, Adeps Lanæ, Cera Flava and Sandarach, of each, 2, powdered Orris Root, 9, and Ether, 16

COLLODIUM SALICYLICUM — Salicylic Acid, 60 grains, Flexible Collodion, 1 oz — *Guy's* and other Hospital Pharmacopœias

Salicylic Acid, 12, Acetone Collodion, q s to produce 100 — *B P C*

The *B P C Supplement* now dissolves the Acid in 30 of Acetone before making up to 100 with the Collodion

Salicylic Acid, 1, Flexible Collodion (by weight), 9 — *F'*

COLLODIUM CUM ACIDO SALICYLICO — Salicylic Acid, 20, Spirit of Ether, 20, Collodum, 60 — *Dutch*

COLLODIUM SALICYLICUM COMPOSITUM — Salicylic Acid, 60 grains, Extract of Indian Hemp, 10 grains, Flexible Collodion, to 1 oz

Collodium Salicylicum Compositum *Syn* **COLLODIUM CALLOSUM** — Salicylic Acid, 12, Extract of Indian Hemp, 2, Acetone Collodion, q s to produce 100 — *B P C*

The *B P C Supplement* now dissolves the Acid and Extract in 30 of Acetone before making up to 100 with the Collodion

COLLODIUM LACTO-SALICYLICUM — Salicylic Acid, 60 grains; Lactic Acid, 1 fl drim, Flexible Collodion, to 1 fl oz

COLLODIUM SALICYLICUM CUM ZINCI CHLORIDO Salicylic Acid, 60 grains, Zinc Chloride, 30 grains, Flexible Collodion, 1 oz — *Guy's*

SALICYLIC DRESSINGS — Gauze, Lint, and Wool, 1 p c, Jute, 4 and 10 p c Jap has Cotton 5 p c and Gauze

GLYCERINUM ACIDI SALICYLICI — Salicylic Acid, 1 part, Glycerin, 9 parts Also called Pasta Acidi Salicylici

LOTIO ACIDI SALICYLICI CUM BORACE — Salicylic Acid, 120 grains, Borax, 60 grains, Glycerin, 1 oz, Rectified Spirit, 1 oz, Distilled Water, to 10 oz — *Middlest*

PULVIS SALICYLICUS CUM TALCO (*Dan, Dutch, Ger, Jap, Norw and Swed*) — Salicylic Acid, 3, Wheat Starch, 10, Talc, 87, mix to a fine powder

Pulvis pro pedibus (*Swiss*) — Same formula as above Used in the German Army as a preventive against perspiring and sore feet It is applied dry, on a march daily, or in garrison every two or three days

Pulvis Talci Salicylicus — Salicylic Acid, 3, Boric Acid, 10, Talc in fine powder, 87 — *US N F*

This has been incorporated in the *B P C* under the title **Pulvis Acidi Salicylici Compositus**

SALICYLIC AND CREOSOTE PLASTER MULLS (*Unna*) — Contain $\frac{1}{2}$ grain of Salicylic Acid and 1 grain of Creosote to the square inch, also twice this strength Possess a solvent power on horny epidermis, the Creosote acting as an anæsthetic Also used in the treatment of lupus — *L*, '86, n 574, *B M J*, '87, n 451

Salicylic Acid and Creosote can also be applied as an ointment with Lard and Wax

SEBUM SALICYLATUM — Salicylic Acid, 2, Benzoin, 10, Mutton Fat, 98 — *Austri*

Used in the German Army for sweaty feet and soreness from riding

Salicylic Acid, 2, Benzoic Acid, 1, Prepared Suet, 97 — *Ger*

The German formula appears in the *B P C* as **Sevum Salicylatum**.

UNGUENTUM ACIDI SALICYLICI — Salicylic Acid, 30 grains, Benzoinated Lard, 1 oz, melt over a water-bath, and stir till cold

Used for eczema, psoriasis, ringworm, and for foul ulcers

VASOLIMENTUM SALICYLICUM — Salicylic Acid, 2, Liquid Vasoliment, 98 — *Huger*

Parogen Salicylatum. *Syn* SALICYLATED VASOLIMENT—Salicylic Acid, 10, Parogen, q s to produce 100—*B.P.C.*

SALACETOL Is obtained by the action of Monochloro-acetone on Sodium Salicylate. Crystallises in long needles, melting at 71°C (159.8°F), soluble 1 in 2,200 of Water, 1 in 15 of Alcohol. It is unaffected by dilute acids, but decomposed by weak alkali with liberation of Salicylic Acid. Introduced as an intestinal disinfectant, resembling Salol in its action—*B.M.J.E.* '96, i 92; *L.* '96, ii 1821

Dose. 15 to 30 grains = 1 to 2 grammes, for adults, 4 to 8 grains = 0.26 to 0.52 gramme for children

ACETYSALICYLIC ACID Aspirin $\text{C}_9\text{H}_8\text{O}_4$, $\text{C}_9\text{H}_7\text{O}_4$, eq. 178.71
Minute, white, odourless crystalline needles. Soluble 1 in 100 Water, 1 in 5 of Alcohol (90 p.c.), soluble in Ether.

Dose. 10 to 15 grains = 0.65 to 1 gramme, three times a day

Antipyretic and antirheumatic—Given as a substitute for the Salicylates in rheumatism, also in pleurisy, the advantage being that it does not produce gastric irritation nor ringing in the ears to the same extent as the Salicylates. Is best prescribed in the form of tablets or wafers—*B.M.J.E.* '99, ii 3, 68, 96, '01, i 92; *i.* 56, *L.* '99, ii 219, *P.J.* '99, ii 135, '00, ii 731, 775, '01, i 665

In various children's diseases, rheumatic affections, and in whooping cough. Action slower than Sodium Salicylate, but effect is more prolonged—*P.* lxx 111, *B.M.J.E.* '02, i 12

15 grain doses once or twice daily given with beneficial effects in cases of inoperable carcinoma—*L.* '03, i 984

In chorea, 10 to 15 grains 3 or 4 times a day—*L.* '03, i 526

Acetylsalicylic Acid should not be prescribed with alkalis or Sodium Bicarbonate—*P.J.* '03, i 2, 89

Superiority of Aceto-salicylic Acid in rheumatic fever, in certain cases when a salicylate has had a good trial—*L.* '07, i 783

In diabetes, alkalis (Soda Water, etc.) and Milk should not be taken with, or just after, the Aspirin—*P.* '07, i 139

Is stated (*L.* '05, i 81) not to possess any advantage over Salicylic Acid and the Salicylates. In some patients dyspeptic symptoms followed its use, in addition to very profuse perspiration

This drug still continues to be very largely used, and numerous references to its advantages appear in Continental literature. Two cases are recorded (*B.M.J.* '05, ii 21) in which a dose of 15 grains caused violent palpitation, difficult respiration, a feeling of extreme weakness and gradually upon lying unconsciousness with voiding of dark green urine. On continuing with 4 grain doses good results were obtained. Toxic symptoms following the administration of Aspirin (*B.M.J.* '06, ii 1692), 100 grains having been taken in all, in 10 grain doses. The acute inflammation of the middle ear attributed to the Aspirin indicates caution in the administration of the drug in any cases complicated with ear trouble. $7\frac{1}{2}$ grains suggested as a sufficiently large dose to begin with. If no unpleasant results occur after taking this amount, it can be then easily increased.

Foreign Pharmacopœias. Official in Dan., Jap. and Swiss, Vendum Acetyl Salicylicum

Tests Aspirin melts at 135°C . (275°F), when warmed with Potassium Hydroxide solution it is saponified with the formation of Potassium Acetate and Potassium Salicylate. The cooled solution when acidified with diluted Sulphuric Acid yields a crystalline precipitate of Salicylic Acid, which, when removed, carefully washed till free from mineral acid and dried, possesses the melting point and answers the test characteristic of Salicylic Acid. A portion of the filtrate when warmed with concentrated Sulphuric Acid and a little Alcohol (90 p.c.) evolves a characteristic odour of Acetic Ether. A weighed quantity of 0.1 of a gramme of Aspirin treated with 5 c.c. of Alcohol and diluted with 20 c.c. of Water should not be coloured violet on the addition of 1 drop of diluted Ferric Chloride T.S., indicating the absence of free Salicylic Acid. A weighed quantity of 0.5 of a gramme should leave no weighable residue when ignited with free access of air, indicating the absence of mineral residue.

Novaspirin, a new preparation of Aspirin, it is said to produce no unpleasant gastric symptoms. It is not so strong, and therefore can be given for longer periods and in larger doses — *B M J E* '07, i 79

AGATHIN — A compound of Salicylic Aldehyde with Methyl phenylhydrazine. Pale greenish crystals, insoluble in Water, soluble in Alcohol (90 p c) and Ether. Has been recommended as an analgesic in sciatica, rheumatic and neuralgic affections — *M A* '95, 8, 608, *Y B T* '94, 463, unreliable and dangerous — *B M J* '98, ii 1055

Dose — 5 to 10 grains = 0.32 to 0.65 grammes

GLYCOSAL (Monosalicylic Acid Glycerin Ester) — White crystalline powder, moderately soluble in Water, readily in Alcohol (90 p c). Introduced as a substitute for Salicylates. Antiseptic, antirheumatic.

Dose, — 5 to 30 grains = 0.32 to 2 grammes

SALITANNAL — A condensation product of Salicylic Acid and Gallic Acid. Introduced as an antiseptic application for wounds.

Iodo-Salicylic Acid and **Di-Iodo Salicylic Acid** are Iodine compounds of Salicylic Acid in which one or two atoms of Hydrogen respectively are replaced by Iodine — *B M J* '97, ii 734

ACIDUM SULPHURICUM.

SULPHURIC ACID

FR, ACIDE SULFURIQUE OFFICINAL, GER, SCHWEFELSAURE, ITAL, ACIDO SOLFORICO, SPAN, ACIDO SULFURICO

A heavy, colourless, odourless liquid, of oily consistence, possessing a strong corrosive action. It may be produced by the suitable oxidation of Sulphurous Anhydride, itself a product of the oxidation of Sulphur or of the combustion of pyrites. It is officially required to contain about 98 p c by weight of Hydrogen Sulphate, H_2SO_4 , eq 97.34

A fuming Sulphuric Acid is known under the title of **Nordhausen Sulphuric Acid**, and is prepared by the distillation of dry Ferrous Sulphate.

Under the name of **Solid Sulphuric Acid**, Sulphuric Anhydride has been introduced into commerce.

Medicinal Properties.—A powerful caustic, and when so used it is made into a paste with an equal quantity of charcoal. In the form of Nordhausen Sulphuric Acid it has been used in cancer (*see Michel's Paste*, p 83). When diluted it is a tonic refrigerant, exciting the appetite and promoting digestion, it is a valuable intestinal astringent, and therefore it is useful in controlling diarrhoea, it diminishes night sweating, more particularly when given with Zinc Sulphate, useful in treating chronic lead poisoning, given with doubtful success in hæmatemesis, hæmaturia and hæmoptysis.

Incompatibles — Alkalis and their Carbonates, salts of Calcium and Lead.

Official Preparations.—Acidum Sulphuricum Aromaticum, and Acidum Sulphuricum Dilutum. Used in the preparation of Acidum Hydrochloricum, Acidum Nitricum, Acidum Sulphurosum, Aether, Aether Aceticus, Cupri Sulphas, Ferri Sulphas, Liquor Ferri Persulphatis, Magnesi Sulphas, Potassii Sulphas, Sodii Sulphas, Sodii Sulphocarbolas, Spiritus Aetheris Compositus, Spiritus Aetheris Nitrosi, Zinci Sulphas and Zinci Sulphocarbolas. **Aromatic**

Sulphuric Acid is contained in Infusum Cinchonæ Acidum. **Dilute Sulphuric Acid** is obtained in Infusum Posa Acidum. Used in the preparation of A. 1. 1. 1. Hydræ, A. 1. 1. 1. Dilutum, Antimonium Sulphuratum and Atropine Sulphas.

Not Official. Tincture Acidæ, Hæmorrh. Mistura Acidæ Sulphuricæ Aromatica, Mixtura Chlorici, Vinum Acidæ, Vinum Chlorici.

Antidotes. In case of poisoning by Sulphuric Acid Magnesia is preferable to chalk. For other antidotes see Hydrochloric and Nitric Acids.

Foreign Pharmacopœias. Official in all the Pharmacopœias ranging from sp. gr. 1.83 to 1.84. *U.S.P.* sp. gr. not below 1.826 at 25° C (77° F). *Fr.* 1.84. *Ger.* 1.84. *Swed.* and *Swiss* contain also a crude Acid.

Tests. Sulphuric Acid has a specific gravity of 1.843, the *U.S.P.* states not below 1.826 at 25° C (77° F), the *P.G.*, 1.836 to 1.840. neither the *R.P.* nor the *P.G.* gives a boiling point for the acid the *U.S.P.* states 338° C (640.4° F). The distinguishing test is the production of a white precipitate, insoluble in Hydrochloric Acid, when Barium Chloride Solution is added to its diluted or neutralised aqueous solutions. It is officially required to contain 97.82 p.c. w/w of Hydrogen Sulphate as volumetrically determined by suitably diluting with Water and titrating with Volumetric Sodium Hydroxide Solution, each gramme should require 20.1 c.c., the *U.S.P.* should contain not less than 92.5 p.c., and *P.G.* 91 to 98 p.c. of absolute Acid.

The more generally occurring impurities are Arsenic and Lead, Selenium, Nitrous, Nitric and Sulphurous Acids. Unlike the *U.S.P.* and *P.G.*, the *R.P.* indicates no special test for either Arsenic or Lead, but is content to group these two important impurities with others of considerably less importance under the elastic expression 'it should yield no characteristic reaction,' etc. The *U.S.P.* employs the modified Gutzeit's test for Arsenic, whilst the *P.G.* employs the Bettendorfs test with Stannous Chloride Solution, both are given in the small type below. A standard of 5 parts per 1,000,000 for Arsenic by weight and 20 parts per 1,000,000 for Lead has been suggested — *C.D.* 08, 1795. *U.S.P.*, in their test for Lead, have adopted a time limit within which no precipitation shall occur when the Acid is carefully mixed with 4 or 5 volumes of Alcohol (91.9 p.c.). It is required to leave no appreciable residue on evaporation.

Selenium, Nitrous, Nitric and Sulphurous Acids may be detected by the tests given in the small type below under the headings of Hydrochloric Acid and Sodium Sulphite, Ferrous Sulphate and Sulphuric Acid, Potassium Permanganate Solution.

Ferrous Sulphate and Sulphuric Acid. If Ferrous Sulphate T.S. be carefully poured as a layer on Sulphuric Acid in a test tube, there should be no coloured zone at the junction of the liquids. *P.G.* and *U.S.P.*

Sulphuric Acid diluted with 20 volumes of Water should not respond to the following tests. It should yield no precipitate, nor become turbid with T.S. of Silver Nitrate, *P.G.* and *U.S.P.* It should not respond to the time limit test for heavy metals, *U.S.P.* and when nearly neutralised with Solution of Ammonia should not be affected by Hydrogen Sulphide Solution. *P.G.* When supersaturated with Ammonia Water, evaporated and ignited, no appreciable fixed residue should remain. *U.S.P.*

Potassium Permanganate Solution.—Sulphuric Acid diluted and should not, on the addition of Potassium Permanganate Solution, imme-

diately discharge its colour *PG* gives 10 c c of Acid diluted with 5 volumes of Water and 3 or 4 drops Potassium Permanganate Solution, *USP* gives 1 c c of Acid, 5 c c of Water, and 0 1 c c Potassium Permanganate Solution

Modified Gutzeit's Test—5 c c of a 1-10 dilution of Sulphuric Acid should not respond to the modified Gutzeit's test for Arsenic, *USP*

Stannous Chloride—No dark colour should be produced in the course of an hour in a mixture of cooled diluted Sulphuric Acid (1-3) and 3 c c Stannous Chloride Solution, *PG*

Hydrochloric Acid with Sodium Sulphite—Let Hydrochloric Acid containing Sodium Sulphite be carefully poured on an equal volume of Sulphuric Acid contained in a test-tube. At the junction of the liquids there should be no red coloured zone, and on warming no reddish precipitate should be formed, *BP*, *PG*, and *USP* (for Selenium), *PG* and *USP* give quantities, viz, 2 c c of each Acid and a fragment of Sodium Sulphite is dissolved in the Hydrochloric Acid

Volumetric Determination—3 c c of Sulphuric Acid are accurately weighed and diluted with 50 c c of Water. The solution is then titrated with Normal Volumetric Potassium Hydroxide Solution, using Methyl Orange Test solution as indicator. The number of c c of alkali used is multiplied by 4 8675 and divided by the weight of Acid taken, the quotient representing the percentage of absolute Sulphuric Acid present, *USP*

Preparations

ACIDUM SULPHURICUM AROMATICUM. AROMATIC SULPHURIC ACID *BP Syn*—ELIXIR OF VITRIOL

Mix gradually 6 of Sulphuric Acid with 59 of Alcohol (90 p c), add Tincture of Ginger 20, and Spirit of Cinnamon 1

Tests.—The specific gravity should be 0 922 to 0 926, and 1 gramme should require 2 84 c c of the Volumetric Sodium Hydroxide Solution for neutralisation, indicating the equivalent of 13 8 p c w/w of absolute Sulphuric Acid. No reference to an indicator of neutrality is given. The *USP* requires a specific gravity at 25° C (77° F) of about 0 933, and that when suitably titrated with Volumetric Solution of Potassium Hydroxide, with Methyl Orange Test-solution as indicator, it shall not indicate less than 20 p c of absolute Sulphuric Acid by weight partly in the form of Ethyl Sulphuric Acid

Dose—5 to 20 minims = 0 3 to 1 2 c c

Sulphovinic Acid stated to occur in Acidum Sulphuricum Aromaticum, its quantity being dependent on age of sample - *PJ '02*, n 137, *CD '02*, n 292

Foreign Pharmacopœias—Dutch and Jap (Tinctura Acida Aromatica), Cort Cinnamon 5, Rad Zingib 5, Acid Sulphurici 10, Spiritus Diluti 90. Mex (Acido Sulfurico Aromatico), Sulphuric Acid 10, Tincture of Ginger 5, Tincture of Cinnamon 5, Alcohol 80. *US* (Acidum Sulphuricum Aromaticum), Sulphuric Acid 111, Tincture of Ginger 50, Oil of Cinnamon 1, Alcohol sufficient to measure 1000, add the Sulphuric Acid gradually and with great caution to 700 of Alcohol and allow it to cool, then add to it the Tincture of Ginger and Oil of Cinnamon, and finally enough Alcohol to make the product measure 1000. Tinctura Aromatica Acida, Norw, Acid Sulph 1½, Aromatic Tincture 8½, Swed, Acid Sulph 1, Aromatic Tincture 19. Not in Ger or Russ.

Tinctura⁴ Aromatica (Dan, Norw and Swed) Cardamoms, 1, Cloves, 1, Galangal Root, 1, Ginger, 1, Cinnamon, 4, diluted Alcohol, 40

Ger, Russ and Swiss contain same ingredients, but have Ginger, 2, Cinnamon, 5, diluted Alcohol, 50

Austr., Cinnamon, 5, Ginger, 2, Zedoary, 1, Caryophyllus, 1, Cardamom, 1; Alcohol (68 p c), 50. It should not yield less than 1.5 p c of solid residue. Jap., Cloves, 2, Cinnamon, 10; Cardamom, 2, Ginger, 5, Dilute Alcohol, 100, extract in the cold for seven days, press, filter, and to the filtrate add Spirit of Lemon, 5.

U.S.N.F. preparation is practically identical with that official in *P.G.*

See also below *Liquor Acidus Halleri*

ACIDUM SULPHURICUM DILUTUM. DILUTED SULPHURIC ACID

Mix gradually 4 of Sulphuric Acid with 40 of Distilled Water, and when cold add more Distilled Water to make 48½ of Dilute Acid at 60° F.

A clear, colourless liquid, possessing a strong acid reaction

As great heat is developed in mixing strong Sulphuric Acid and Water, it is always safer to add the Acid to the Water than the Water to the Acid. When Acid 1, Water 1, were mixed the temperature rose to 270° F (132.2° C.)

12 minims contain about 1 minim of strong Sulphuric Acid

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

Prescribing Notes. *Prescribed, much diluted, in mixtures; or in cough liniments, with Squill, Poppies, and Syrup of Mulberry, also to dissolve Quinine*

Foreign Pharmacopœias.—Official in Austr., Acid 1, Water 4 76, sp. gr. 1.12; Ital., Acid 1, Water 4, sp. gr. 1.180 Dutch, sp. gr. 1.124, Ger. and Russ., Acid 1, Water 6, sp. gr. 1.110 to 1.114, Dan. and Norw., Acid 1, Water 7, sp. gr. 1.081 to 1.085, Belg., 1:1; Hung., Jap. Port. and Span., Acid 1, Water 9; Fr., Swed., Swiss, 10 p c, U.S., 10 p c, sp. gr. about 1.067 at 25° C. (77° F.). All by weight.

Tests.—Diluted Sulphuric Acid is officially required to possess a specific gravity of 1.094, and to contain 13.63 p c of Hydrogen Acetate, as officially determined by titration with Volumetric Solution of Sodium Hydroxide, the *U.S.P.* diluted acid is required to have a specific gravity of about 1.067 at 25° C. (77° F.), and to contain not less than 10 p c w/w of absolute Sulphuric Acid as determined by titration with Normal Volumetric Potassium Hydroxide Solution, using Methyl Orange Test-solution as an indicator, that of the *P.G.* has a specific gravity of 1.110 to 1.114, and contains from 15.6 to 16.3 p c. w/w of pure acid

The Diluted Acid should not fail to respond to the tests of identity and purity given under *ACIDUM SULPHURICUM*, but *B.P.* does not say so, the *U.S.P.*, on the other hand, duly notes this.

Not Official.

LIQUOR ACIDUS HALLERI. *Syn.* ACIDUM SULPHURICUM ALCOHOLICUM, MISTURA SULPHURICA ACIDA, AQUA RABELLI, AQUA RABELII, AQUA DE RABEL, EAU DE RABEL.

Austr., Ger., Hung., Mex., Port., Russ., Span. and Swiss.—Sulphuric Acid, 1, Alcohol (90 p c), 3

Dan., Dilute Sulphuric Acid, 1; Syrup of Raspberries, 9; Distilled Water, 40.

Fr., Sulphuric Acid, 1, Alcohol (95 p c), 3; Poppy Petals, 0.04.

Dutch, Ital. and Norw.—Sulphuric Acid, 1; Alcohol, 1.

All by weight.

MISTURA ACIDI SULPHURICI AROMATICA.—Aromatic Sulphuric Acid, 10 minims, Red Mixture, to 1 oz.

MYNSICHT'S ELIXIR OF VITRIOL.—Cinnamon, Ginger and Cloves, of each 3, Calamus Aromaticus, 8, Galangal, 12, Sage, 4, Peppermint, 4, Cubebs, 2, Nutmeg, 2, Aloes Wood, 1, Lemon Peel, 1, Sugar Candy, 32, Alcohol (90 p c), by weight, 144, Sulphuric Acid, by weight, 96. Digest for three weeks.

Dose.—5 to 10 minims = 0.3 to 0.6 gramme

MICHEL'S PASTE.—Nordhausen Sulphuric Acid, 3, by weight, Asbestos, finely powdered, 1. Should be prepared fresh as required.

ACIDUM SULPHUROSUM.

SULPHUROUS ACID

FR, ACIDE SULFUREUX, GER, SCHWELFIGSAURE, ITAL, ACIDO SOLFOROSO

A colourless liquid, with a strong characteristic odour of burning Sulphur, officially required to contain 6.4 p c of Hydrogen Sulphite, H_2SO_3 , eq 81.46

It is prepared by the oxidation of Sulphur or by the reduction of Sulphuric Acid by boiling with Carbon or Copper.

Medicinal Properties.—It is a powerful deoxidising agent, disinfectant and antiseptic. In 1 dram doses, freely diluted, it is valuable in vomiting depending on fermentation in the stomach, and as an intestinal antiseptic in enteric fever. Diluted with 1 or 2 parts of Water, it is used as a **spray** in diphtheria and follicular tonsillitis, mixed with equal parts of Glycerin, as an **application** in erysipelas, ringworm, and other parasitic skin diseases. Also for chapped hands and chilblains, very effectual in **chapped nipples**, as a **lotion**, 1 or 2 drms to 1 oz of Water, for **verrucae**, cuts, ulcers, and bed-sores, as an **inhalation** in nasal catarrh and influenza, 60 minims in 20 oz of Water at 60° to 100° F.

Pfeiffer found that 0.5 to 1 p c aqueous solution caused excessive and extensive gastritis. Even 20 minims largely diluted caused irritation of the digestive organs (*A J P*, '90, 626). Brunton, however, strongly recommends 1 dram doses thoroughly diluted, in gastric fermentation, 20 to 30 minims every two or three hours, stated (*B M J* '04, 11 1450) to check fermentative changes in the bowel in enteric fever.

Dose.— $\frac{1}{2}$ to 1 fl dram = 1.8 to 3.6 c c

Foreign Pharmacopœias.—Official in Port, Soluto de Gaz Sulfuroso, U S, 6.0 p c, SO_2 .

Tests.—Sulphurous Acid has a specific gravity of 1.025; the *U S P* gives not less than 1.028 at 25° C (77° F), it is not official in the *P G*. The distinguishing tests are its strong sulphurous odour, its power of decolorising Iodine Solution (which is utilised in its quantitative determination), the production of Hydrogen Sulphide when reduced by Hydrochloric Acid and Zinc, and its bleaching action on certain vegetable colouring matters.

It is officially required to contain 6.4 p c of Hydrogen Sulphite, as determined by titration with Volumetric Solution of Iodine, using Starch Mucilage as an indicator.

The most likely impurities are an undue amount of Sulphate due to oxidation on keeping and the presence of other Sulphur compounds,

notably Hydrogen Sulphide and mineral residue. The presence of Sulphate is guarded against by tests with Barium Chloride Solution, and the mineral residue by the fact that it is required to leave no residue on evaporation.

Mercurous Nitrate Solution—The gas, evolved on gently heating a few c.c. of the Acid in a test-tube, blackens a strip of paper moistened with T.S. of Mercurous Nitrate, *U.S.P.*

Lead Acetate Solution.—The gas, evolved on gently heating a few c.c. of the Acid in a test tube, does not affect a strip of paper moistened with T.S. of Lead Acetate, *U.S.P.*

Barium Chloride Solution. It is officially required to yield only a slight precipitate with Barium Chloride Solution, but if the Sulphurous Acid be oxidised by the cautious addition of Chlorine Solution it yields on the addition of Barium Chloride Solution a copious white precipitate. The *U.S.P.* test gives quantities, 1 c.c. Sulphurous Acid diluted with 99 c.c. of Water, 1 c.c. diluted Hydrochloric Acid, and then 1 c.c. Barium Chloride T.S. yielding not more than a slight turbidity at once.

Volumetric Determination. 1 gramme of the Acid diluted with about 100 c.c. of Water requires 15 c.c. of Volumetric Iodine Solution, *B.P.* 2 c.c. of the Acid are accurately weighed in a stoppered weighing bottle. To this are added 50 c.c. of Decinormal Volumetric Iodine Solution, and the mixture is allowed to stand for 5 minutes. Tenth normal Volumetric Sodium Thiosulphate Solution is now slowly added until the mixture is just decolorised. Subtract the number of c.c. of the Tenth-normal Volumetric Sodium Thiosulphate Solution used from 50, and multiply the difference by 0.118, and divide this product by the weight of the Acid taken, the quotient represents the percentage of absolute Sulphurous Acid in the latter, *U.S.P.*

Not Official.

SULPHUR DIOXIDE LIQUEFIED—This is also commonly known as **Liquefied Sulphurous Acid Gas**. It is supplied in siphons, and in tinned iron vessels with soft lead exit tube.

Disinfecting with Sulphur.—This is usually done with liquid Sulphur Dioxide. The room to be disinfected should be sealed up so as to prevent any ventilation, by blocking up the fireplace and pasting paper over the cracks of the windows. The small leaden exit-tube of the vessel is cut in the room, so as to allow the gas to escape somewhat slowly, and the operator retires quickly, shuts the door and papers up the cracks of the door so as to complete the sealing of the room. The room is allowed to remain closed for 12 hours, and then opened cautiously. About 20 oz. gas is required for a room of a size 1700 cubic feet. When the liquefied gas is unobtainable, Sulphur which is sold in the form of Sulphur Candles can be used as a substitute, but the gas is much more effectual.

In order to obtain the maximum disinfecting power of the Sulphur Dioxide it is necessary to introduce moisture, and this may be done by placing open pans of steaming water in the room, or by injecting steam.

ACIDUM TANNICUM.

TANNIC ACID.

B.P.Syn.—TANNIN.

$C_{14}H_{10}O_9$, eq. 319.66

Fr., TANNIN OFFICINAL, GER., GEBEHALLE, ITAL., ACIDO TANNICO; SPAN., ACIDO TANICO.

B.P. gives the formula as $C_{14}H_{10}O_9$, $2H_2O$, but the $2H_2O$ is discountenanced by most of the standard works on chemistry.

A pale buff-coloured, micro-crystalline powder, possessing an acid reaction and a characteristic astringent taste. It is obtained from Nut Galls.

Solubility.—10 in 5 of Water, 10 in 6 of Alcohol (90 p.c.), 3 in 1 of Absolute Alcohol, 1 in 3 of Glycerin, or if warmed, 1 in 1, sparingly in Olive Oil, almost insoluble in Benzol, in Chloroform, and in Ether.

These solubilities were made with Tannic Acid which was very soluble, but different samples vary in solubility.

Commercial Tannic Acid frequently contains some proportion of Gallic Acid, which when dissolving in Water is the last portion to go into solution, and which may be detected by the Potassium Cyanide test mentioned under Gallic Acid.

Medicinal Properties.—Styptic and local astringent. 60 grains in 10 oz. of Rose Water are used as a **spray** for relaxed sore throat, the same strength is also used as an **injection** in leucorrhœa and in chronic gonorrhœa with advantage, 3 grains to the oz. is used as a **nasal douche**, 60 grains to the oz. as an **ointment**, the powder has been used as a **snuff** in epistaxis. Internally for gastric and intestinal hæmorrhage acting as a direct styptic. A dose of 1 drim. is often successful in hæmorrhage from gastric ulcer. For **suppositories** and **pessaries**, see p. 87. The **glycerin** is used as a **paint** in relaxed throat, and for nasal discharges, also locally as a styptic.

Equal parts of Glycerin of Tannin and Glycerin of Alum form a good application for relaxed throat.

As an injection into nasal polypi — *L* '87, i. 543.

Warm Tannin enemata were given with success in the cholera at Naples — *L* '85, i. 352.

Tannic Acid is no doubt a local styptic by its albumen-coagulating power. But the careful researches of Stockman have demonstrated the futility of using it as a remote astringent — *B M J* '00, ii. 1070.

A solution, 2 in 10 of spirit, the best application in alveolar pyorrhœa, loose teeth becoming tight, and regaining their power of mastication, also in dental neuralgia, applied to the gums and round the teeth — *B M J E* '07, ii. 12.

Dose.—2 to 5 grains = 0.13 to 0.32 gramme.

Prescribing Notes.—*Prescribed in Water, and may be combined with the Ferrous (but not with the Ferric) salts of Iron. Can be given in cachets or Compressed Tablets. 4 grains with ½ minim of Glycerin make a nice pill. 60 grains to 1 oz. of Chalk with 30 grains of Powdered Soap make an astringent dentifrice.*

Incompatibles.—Mineral Acids, Alkalis, Antimony salts, Lead and Silver, Ferric salts, the vegetable alkaloids, and Gelatin.

Official Preparations.—Glycerinum Acidî Tannici, Suppositoria Acidî Tannici, and Trochiscus Acidî Tannici.

Not Official.—Crayons de Tannin, Gargarisma Acidî Tannici, Lotic Acidî Tannici, Sulphureosa, Nebula Acidî Tannici, Pessary or Vaginal Suppository, Schuster's Pastilles, Suppositorium Acidî Tannici c. Opio, Supp. Ac. Tann. et Belladonnæ, Supp. Ac. Tann. et Morphine, Unguentum Acidî Tannici, Unguentum Acidî Tannici c. Opio, Tannic Wool, Tannalbin, Honthin, Glutanol, Tannigen, Tannoform, Tanocol, and Tannone.

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests.—The distinguishing test for Tannic Acid is its behaviour with Ferric Chloride Test-solution, which yields a bluish black

colour or precipitate. The *U.S.P.* states that a 1 p.c. w/v aqueous solution of Tannic Acid yields on the addition of a small quantity of Calcium Hydroxide Test-solution a pale bluish-white flocculent precipitate which is not dissolved on shaking, and which becomes more copious and of a deeper blue on the addition of a moderate excess of Calcium Hydroxide Test-solution, a large excess imparting a pale pinkish tint.

Tannic Acid may be distinguished from Gallic Acid by its aqueous solution yielding precipitates with solutions of Isinglass, Albumen or alkaloids. The *U.S.P.* states that it may be distinguished by the test with Calcium Hydroxide Test-solution given above, and by the fact that its aqueous solution produces precipitates with most alkaloids and glucosides, and with Test-solutions of Gelatin, Albumen and Starch. The *B.P.* states Tartarated Antimony, but it has been shown that the latter reagent produces a precipitate in Gallic Acid Solutions also; the *U.S.P.* omits reference to Tartarated Antimony.

Mineral acids and certain mineral salts precipitate Tannic Acid from solution, and the filtrates obtained possess nostringency; the *P.G.* states that it is precipitated from its 1 to 5 aqueous solution by the addition of Sulphuric Acid or by Sodium Chloride.

The most likely impurities are Gum or Dextrin, resinous substances and mineral matters. Traces of Gallic Acid may also be present. Both *U.S.P.* and *P.G.* include specific tests for Gum or Dextrin and resinous substances, which are described in the small type below under the heading of Alcohol and Ether, or Alcohol. Gallic Acid may be detected by the Potassium Cyanide test mentioned under Gallic Acid. No appreciable residue should be left on ignition with free access of air. *U.S.P.* requires it to leave not more than 0.2 p.c., and *P.G.* that 0.5 gramme should not leave a weighable residue. Austr. and *P.G.* require that the Acid shall not lose more than 12 p.c. by weight on drying at 100° C (212° F), indicating two molecules of Water of crystallisation.

Alcohol and Ether, or Alcohol. A mixture of 2 c.c. of an aqueous solution of the Acid and 2 c.c. of Alcohol (90 p.c.) should remain clear, also on the further addition of 1 c.c. of Ether, *P.G.* The *U.S.P.* dissolves 2 grammes in 10 c.c. of boiling Water, 5 c.c. of which solution when cooled and diluted with 10 c.c. of Alcohol (94.9 p.c.) should produce no turbidity; or (for resinous substances) when diluted with 10 c.c. of Water.

Preparations.

GLYCERINUM ACIDI TANNICI. GLYCERIN OF TANNIC ACID.

Tannic Acid, 1; Glycerin, sufficient to produce 5. (1 in 5)

Foreign Pharmacopœias - Official in Belg., 8 and 17; Port., 1 and 9; U.S., 1 and 4; Fr. and Mex., 1 and 5 of Glycerin of Starch. Not in the others.

SUPPOSITORIA ACIDI TANNICI. TANNIC ACID SUPPOSITORIES

Contains 3 grains = 0.2 gramme of Tannic Acid in each suppository, mixed with Oil of Theobroma.

TROCHISCUS ACIDI TANNICI. TANNIC ACID LOZENGE

$\frac{1}{2}$ grain Tannic Acid in each, with Fruit Basis

Dose —1 to 6 lozenges

Foreign Pharmacopœias—Official in Jap, $\frac{3}{4}$ grain each, U S, about 1 grain each with Sugar and Orange Flower Water

Not Official

CRAYONS DE TANNIN (*F*)—Tannin, 20, Gum Acacia, 1 (both in powder), mix and make into a mass of pilular consistence by means of equal parts Glycerin and Water, then roll into cylinders of the size required

GARGARISMA ACIDI TANNICI—Glycerin of Tannic Acid, 1 fl oz, Water, to 10 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

Glycerin of Tannic Acid, 1 fl drim, Water, to 1 fl oz—*Charing Cross, London and St Bartholomew's*

LOTIO ACIDI TANNICI SULPHUROSA—Glycerin of Tannic Acid, 1, Sulphurous Acid, 1, Distilled Water, to make 8 *St Bartholomew's*

NEBULA ACIDI TANNICI—Glycerin of Tannin, 1 fl drim, Distilled Water, to 1 fl oz—*City of London Chest*

Glycerin of Tannin, 40 minims, Water, to 1 oz—*Throat*

Glycerin of Tannin, 1, Distilled Water, *q s* to produce 10—*B P C*

PESSARY OR VAGINAL SUPPOSITORY—Tannic Acid, 10 grains Stearin, or Oil of Theobroma, sufficient to make 2 drim, for one pessary Used in leucorrhœa

This has been incorporated in the *B P C*

SCHUSTER'S PASTILLES—Tannic Acid, 30 grains, Opium, 1 grain, Glycerin, *q s* to form suitable cylinders for the male urethra

SUPPOSITORIUM ACIDI TANNICI C OPIO Tannic Acid, 3 grains, Powdered Opium, 1 grain, Stearin, or Oil of Theobroma, 11 grains

SUPPOSITORIA ACIDI TANNICI ET BELLADONNÆ—Tannic Acid, 3 grains, Extract of Belladonna (*B P '85*), $\frac{1}{2}$ grain, Oil of Theobroma, to 15 grains—*St Bartholomew's*

SUPPOSITORIA ACIDI TANNICI ET MORPHINÆ—Tannic Acid, 8 grains, Morphine Hydrochloride, $\frac{1}{2}$ grain, Oil of Theobroma, to 15 grains—*Westminster*

UNGUENTUM ACIDI TANNICI—Tannic Acid, 20, Glycerin, by weight, 20, Ointment, 60—*U S P*

This has been incorporated in the *B P C*, but the simple ointment of the *U S P* is different from the *B P C*, which resembles *B P 1885*

UNGUENTUM ACIDI TANNICI C OPIO Tannic Acid, 30 grains, Powdered Opium, 30 grains, Lard, 1 oz

TANNIC WOOL—Dissolve 2 of Tannic Acid in 60 of Water, and with it thoroughly moisten 8 of Absorbent Cotton-Wool, press so as to remove 30 of the fluid, then dry the Wool in a warm chamber When dry remove any discoloured portion. This is sold as **Wool for cigarettes**.

TANNALBIN A light brown, tasteless powder, insoluble in Water A combination of Tannic Acid with albumin, which by a special treatment has been so altered that it is insoluble in the gastric juice, ordinary Albumin Tannate being readily soluble

Tannalbin and Bismuth Subnitrate given early in the morning, as astringents in treatment of catarrhal ulcers of the large intestine—*B M, J B '99, i 59*

It has been introduced as an intestinal astringent.

Adult Dose.—15 grains = 1 gramme, given at intervals of one or two hours

Foreign Pharmacopœias—Official in Austr., Belg., Dutch and Swed. Not in the others.

Tests.—When shaken with Water and filtered, the filtrate affords with Ferric Chloride Test-solution a bluish-black coloration. When boiled with Water, cooled, and filtered, the filtrate gives a precipitate with Albumin Solution. When shaken with solution of Sodium Hydroxide (15 p.c.) the mixture gelatinises. When further heated to the boiling point, cooled, and supersaturated with Hydrochloric Acid, an odour of Sulphuretted Hydrogen is evolved, and a white precipitate is thrown down.

The Austr. Ph. gives the following test: 1 gramme of Tannalbin digested for 4 hours at 10° C. (25° F.) with 0.1 gramme of Pepsin, 50 c.c. of Water and 1.5 gramme of Dilute Hydrochloric Acid leaves a residue which after washing with 30 c.c. of water and drying at 100° C. (212° F.) shall not weigh more than $\frac{1}{2}$ gramme. The Dutch Ph. requires that it shall not yield more than 1 p.c. of ash.

HONTHIN (Albumin Tannate). Greenish-brown tasteless odourless powder. Insoluble in water.

Dose. 10 to 30 grains = 0.65 to 2 grammes, three to five times a day for adults; 5 grains = 0.32 gramme, for children.

GLUTANOL. Combination of Tannic Acid with vegetable fibrin. Similar in action and properties to Tannalbin. Administered in powder form.

Dose. 5 to 15 grains = 0.32 to 1 gramme.

TANNIGEN (Di-acetyl Tannin).—A greyish-white tasteless powder. Practically insoluble in Water, but readily in alkaline solutions. Recommended in diarrhoea, principally of children, but also in that of adults. It passes through the stomach unchanged, but on entering the alkaline intestinal tract it breaks up and acts as an astringent.

Dose.—1 to 3 grains = 0.06 to 0.18 gramme for children, and 5 to 10 grains = 0.32 to 0.65 gramme for adults. Small doses can be mixed with an equal quantity of Milk Sugar, and larger doses for adults can be taken in *cachets*.

Tests. When shaken with Water and filtered, the filtrate is coloured bluish-black by Ferric Chloride Test-solution. When warmed with Potassium Hydroxide Solution, cooled, mixed with diluted Sulphuric Acid and a little Alcohol (90 p.c.) and warmed, the odour of Acetic Ether is evolved. It should leave not more than 1 p.c. of mineral residue when ignited at a dull red heat.

TANNOFORM (Methyl Ditannin).—A light, pinkish-brown, odourless and tasteless powder. A condensation product of Tannic Acid and Formic Aldehyde. It is claimed to possess the astringent effects of Tannin with the antiseptic and drying properties of Formaldehyde. Used as an application in skin diseases, and to wounds, either alone or mixed with Starch, or diluted 1 to 4 with French Chalk, as a dusting powder.

As a 10 p.c. ointment in eczema. —*B.M.J.E.* '99, ii, 48; *M.I.* '00, 182.

Rubbed into the chest for night sweats in phthisis. —*B.M.J.E.* '01, ii, 50.

Foreign Pharmacopœias—Official in Swed. and Swiss.

TANOCOL. A white or nearly white odourless and tasteless powder, a combination of Gelatin and Tannic Acid. Insoluble in Water. Stated to be useful as an intestinal astringent.

Dose.—10 to 15 grains = 0.65 to 1 gramme, for an adult; 5 grains = 0.32 gramme, for children. —*L.* '03, i, 1039.

TANNONE.—A condensation product of Tannic Acid and Hexamethylene tetramine. A light brown tasteless powder, almost insoluble in Water and weak Acids, and dissolves slowly in weak alkalis.

Dose.—15 grains = 1 gramme for adults, children, 8 to 5 grains = 0.52 to 0.32 gramme.

ACIDUM TARTARICUM.

TARTARIC ACID



FR, ACIDE TARTRIQUE, GFR, WEINSAURF, ITAL, ACIDO TARTARICO, SPAN, ACIDO TARTRICO

Colourless and odourless, translucent monoclinic prisms, possessing a strongly acid taste

It is a di-basic acid prepared from Argol or crude Tartar, and is chemically a Di-oxy-succinic or Di-hydroxy-succinic Acid

Solubility.—10 in 8 of Water, and measures 14, 1 in $2\frac{1}{2}$ of Alcohol (90 p.c.), 1 in $4\frac{1}{2}$ of Glycerin, 1 in 40 of Ether, 1 in 5 of Absolute Alcohol, nearly insoluble in Benzol and Chloroform

The solubility of this acid in Ether naturally varies with the amount of Water and Alcohol contained in the Ether. The above figure represents the solubility in *B.P.* Ether (sp. gr. 0.735), but the figure for Ether Purus (sp. gr. 0.720) is 1 in 195

Medicinal Properties —The same as Citric Acid

Dose —5 to 20 grains = 0.32 to 1.3 grammes

Incompatibles —Salts of Potassium, Calcium, Mercury and Lead, Alkaline Carbonates, and the vegetable astringents

Official Preparations —Used in the preparation of Pulvis Sodæ Tartaricæ Effervescent, Sodii Citro-Tartras Effervescent, and the other granular effervescent preparations

Foreign Pharmacopœias —Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Jap., Ital., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests —The distinguishing test for Tartaric Acid is the formation of a white crystalline precipitate on the addition of Potassium Acetate Solution to its aqueous solution, the precipitate being soluble in Ammonium Chloride Solution or in Sodium Hydroxide Solution, also the mirror test with Silver Nitrate Solution in its neutralised solutions. Its aqueous solution is officially required to be dextro-rotatory. It is officially required to contain 99.0 p.c. of Hydrogen Tartrate, as determined by titration with Volumetric Sodium Hydroxide Solution, 1 gramme when dissolved in Water neutralises 13.3 c.c. of Volumetric Sodium Hydroxide Solution. *U.S.P.* requires it to contain not less than 99.5 p.c. of pure Tartaric Acid, the *P.G.* does not mention a requisite percentage, the *U.S.P.* states that Phenol-phthalein Test solution is to be used as an indicator of neutrality. Tartaric Acid may be distinguished from Citric Acid, and its presence detected in the latter by its power of decolorising a weak solution of Potassium Chromate, upon which Citric Acid has no action (Alcohol and other reducing agents must be absent). The Pusch's test (*P.J.* [3] xv 693) with Sulphuric Acid at 100° C (212° F) which easily detects 1 p.c. of Tartaric Acid in Citric Acid. The Resorcinol Sulphuric test (*C.D.* '91, 16) is also a delicate test for Tartaric Acid, but in the presence of a large proportion of Citric Acid the red colour is

rather obscure, and in that case it offers no advantage over Pusch's test

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron and Lead, Oxalates and Sulphates. The most important impurity is Lead.

A standard of 10 parts per 1,000,000 for Lead, and 1 part per 1,000,000 for Arsenic has been proposed (*C.D.* '08, i. 795). Arsenic, Copper, Iron and Lead may be detected, if present, by the Hydrogen Sulphide test, Calcium by the test with Ammonium Oxalate Solution; Oxalates and Sulphates by the tests with Calcium Sulphate Solution and the Barium Chloride or Nitrate test. Each of these tests is given under its respective heading in the small type below.

When ignited with free access of air it should leave not more than 0.05 p.c. of mineral matter. This is the requirement of both the *B.P.* and the *U.S.P.*; the *P.G.* requires that 0.5 of a gramme of the acid should leave no weighable residue. The *B.P.* standard is considered (*C.D.* '08, i. 795) as too severe, and a limit of 0.1 p.c. of ash is suggested.

Melting Point. 135°C (275°F), *U.S.P.* When kept at a temperature of 100°C. (212°F) for some time, the powdered crystals do not suffer a sensible loss of weight, *U.S.P.*, at a temperature above 135°C. (275°F) the acid chars, emitting an odour resembling that of burning sugar, *P.G.* and *U.S.P.*

Potassium Acetate Solution.—A (1-3) aqueous solution yields with Potassium Acetate Solution a crystalline precipitate, soluble in Ammonium Chloride Solution and in Sodium Hydroxide Solution; the solution in Sodium Hydroxide gelatinises on warming, but becomes fluid again on cooling, *P.G.* The *U.S.P.* Potassium Acetate test is made with a (1-2) aqueous solution and a (1-3) Potassium Acetate solution, and the crystalline precipitate obtained is soluble in solutions of the alkalis and mineral acids, but not in Acetic Acid.

Hydrogen Sulphide.—There should be no darkening in colour within 5 minutes in a solution of 10 grammes of Tartaric Acid in 20 c.c. of Water nearly neutralised with solution of Ammonia and sufficient saturated solution of Hydrogen Sulphide added to produce 100 c.c., *B.P.*

It should not, after the addition of a few drops of Hydrochloric Acid, show any colour when submitted to the time-limit test for heavy metals, omitting the subsequent addition of Ammonia Water, *U.S.P.*

5 grammes dissolved in 10 c.c. of Water, and solution of Ammonia added until only faintly acid, should be unaffected by Hydrogen Sulphide Solution, *P.G.*

Barium Chloride or Barium Nitrate.—A (1-10) aqueous solution of Tartaric Acid should be unaffected by Barium Nitrate Solution, *P.G.*, by Barium Chloride T.S. after acidulation with a few drops Hydrochloric Acid, *U.S.P.*

Ammonium Oxalate.—A (1-10) aqueous solution of the acid should be unaffected by Ammonium Oxalate Solution, *P.G.*, *U.S.P.* allows a faint turbidity in 10 c.c. of a solution (1-10) acidulated with Ammonia Water on the addition of Ammonium Oxalate T.S.

Calcium Sulphate Solution.—A (1-10) aqueous dilution should be unaffected by Calcium Sulphate T.S. after nearly but not quite neutralising with Solution of Ammonia, *P.G.* and *U.S.P.*

Volumetric Determination.—3.73 grammes should require not less than 49.8 c.c. normal Potassium Hydroxide Volumetric Solution (each c.c. corresponding to 2 p.c. of pure Tartaric Acid), *U.S.P.* Phenolphthalein Test-solution is used as the indicator.

Not Official.

ACONITI FOLIA.

ACONITE LEAVES

The fresh leaves and flowering tops of *Aconitum Napellus*, L., gathered when about one-third of the flowers are expanded, from plants cultivated in Britain.

This plant and the Extract from the fresh herb were formerly official, but are now omitted.

Foreign Pharmacopœias—Official in Fr, Mex, Port, Russ and Span. Not in the others.

Descriptive Notes.—The leaves only of the plant having flowers shaped like a shallow or navicular helmet should be used, since in some districts *Aconitum paniculatum*, Lam., is cultivated for sale on account of giving a smoother and less resinous extract. It differs from *A. Napellus* chiefly in the helmet being twice as deep as broad and in the more branched inflorescence, and as it does not contain Aconitine, and consequently does not produce the characteristic tingling and numbing sensation when chewed, the leaves should not be used in medicine for that drug. The leaves are most active when about one-third of the flowers are expanded.

Aconite leaves are rarely used in the dried state. The lower leaves are long-stalked and have five or seven lobes, each of which is pinnatifid with linear acute segments. The upper leaves have three to five lobes, they are quite smooth, and paler beneath. The leaves are most active before the flowers are fully formed.

For microscopical detail see *Vogl Anat Atlas Tab 16*.

The dried leaves imported from Germany should not be used since it is impossible to distinguish the active and inactive leaves when mixed, and unless the flowers are present there can be no certainty as to the species collected.

ACONITI RADIX.

ACONITE ROOT

FR, TUBERCULE D'ACONIT, GER, EISENHUTKNOLLEN, ITAL, TUBERO DI ACONITO, SPAN, ACONITO

The root collected in the autumn from *Aconitum Napellus*, L., cultivated in Britain, and dried.

The *USP* orders 'root collected in autumn, containing not less than 0.5 p.c. of Aconitine', *PG* states 'collected at the end of flowering time and requires it to yield not less than 0.5 p.c. of alkaloid reckoned as Aconitine'. *BP*, *USP* and *Ger* all specify the root of *Aconitum Napellus*. *BP* does not give a standard of alkaloid.

Medicinal Properties—Anodyne, antiphlogistic, antipyretic, diaphoretic. Externally it relieves the pain of acute and chronic rheumatism, facial neuralgia, and of itching, as in erythema. Given internally it lessens the frequency and tension of the pulse, relieves pain and high temperature, and is thus useful in all acute local inflammations (not advanced), such as those of pneumonia, eruptive fevers, erysipelas, tonsillitis, peritonitis, and painful neuralgic affections, contra indicated when valvular disease of heart is present, or when fever is prolonged as in typhoid.

Beneficial in lowering blood pressure in acute uræmia.—*B.M.J.* '06, ii. 1450.

It is better given in small doses and very frequently, $\frac{1}{4}$ to 1 minim of Tincture every ten minutes or quarter of an hour for two hours, then hourly.—*Ringer*

Five minims of Tincture given every three or four hours, increasing the dose to 20 minims, succeeded in curing a case of neuralgia in the face, when every other remedy tried had failed

Ph. Ger. maximum single dose, 0.1 gramme; maximum daily dose, 0.3 gramme

Antidotes.—In case of poisoning by Aconite, use emetics, Apomorphine $\frac{1}{16}$ grain, Alcoholic stimulants, Atropine or Belladonna, Digitalis, Amyl Nitrite

Atropine is antagonistic to the action of Aconitine on the heart

Official Preparations.—Lanimentum Aconiti and Tinctura Aconiti Used in the preparation of Aconitina

Not Official.—Extractum Aconiti Radicis Alcoholicum, Chloroformum Aconiti, Lanimentum Aconiti et Chloroformi, Lanimentum Aconiti Compositum, Pastillus Aconiti, Pigmentum Iodi cum Aconito, Tinctura Aconiti Fortior and Trochisci Aconiti

Foreign Pharmacopœias.—Official in Belg., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Port., Russ., Span., Swiss and U.S. Not in the others

Descriptive Notes.—The Aconite root met with in commerce varies considerably in quality and appearance. The root cultivated in Britain is either dried whole or split into two or three longitudinal segments. The latter form enables the soundness of the root to be seen, but the structure is not so easily observed as in the roots dried whole. When collected in autumn the root is solid, brownish externally and white and starchy within; but if collected in summer when in full flower, the old root to which the flowering stem is attached is spongy and porous or hollow, being exhausted by the flowering process, such roots are excluded by the *B.P.* The root dried entire exhibits, when broken across the centre, a seven-angled portion or pith, with a small group of vessels visible at each angle, and well-marked dark cambium line, and is, if carefully dried, white and starchy, but if overheated it has a resinous fracture. It is about $2\frac{1}{2}$ to $3\frac{1}{2}$ in. (62 to 87 mm) long, and $\frac{3}{8}$ in. to $\frac{7}{8}$ in. (15 to 22 mm) in diameter at the widest portion. If cautiously applied to the tongue, it should cause a numbing sensation after a short interval; the *Aconitum pauciflorum*, Lam., does not do so. The root collected in autumn consists of the new root, and may be recognised by having only leaf scales at the apex, and not any portion of stem. The *B.P.* requires the root to be crowned with the remains of an undeveloped bud, and therefore excludes the German root, which is usually collected when the plant is in flower, and consequently crowned with the base of the stem; it is also collected indiscriminately by peasants from any blue-flowered species, and therefore varies considerably in strength. Japanese root is shorter, grey-brown externally, conical, and tapering abruptly to a point; it is obtained from a different species, stated to be *A. Fischeri*, Reicht.

Tests.—Both *U.S.P.* and *P.G.* require the root to yield a definite content of alkaloid; the *U.S.P.* stipulating that it shall yield not less than 0.5 p.c. of Aconitine; the *P.G.* that it shall contain not less than 0.5 p.c. of alkaloid calculated as Aconitine. The process adopted by the *U.S.P.* is essentially as follows.—A weighed quantity of 10 grammes of the root is treated in an Erlenmeyer flask

with 75 c c of a mixture of 7 parts of Alcohol (94.9 p c) and 3 parts of Water, and shaken at intervals for five hours. The contents of the flask are then transferred to a small glass percolator, and after the liquid has passed through the percolation is continued with more of the above mixture of Alcohol and Water until 150 c c of percolate is obtained. The percolate is then evaporated at a temperature not exceeding 60° C (140° F), and 5 c c Tenth-normal Volumetric Sulphuric Acid Solution and 10 c c of Distilled Water added. The liquid is filtered into a separator, the dish and filter being washed with about 40 c c of Distilled Water and the washings added to the separator, 25 c c of Ether and 2 c c Animonin Solution are added, and the mixture agitated for one minute. The lower stratum is drawn off, the Ether-solution is filtered. The lower stratum is returned to the separator and shaken with 15 c c of Ether for one minute, the process of drawing off being repeated. The lower stratum is washed with two further portions of 10 c c each of Ether. The combined Ether-solutions are evaporated to dryness, and the residue dissolved in 3 c c of Tenth-normal Volumetric Sulphuric Acid Solution, and titrated back with Fiftieth normal Volumetric Potassium Hydroxide Solution until a violet colour is produced, 5 drops of Cochineal Test-solution being used as an indicator of neutrality.

The process of the *P G* is essentially as follows.—A weighed quantity of 12 grammes of root, dried at 212° F (100° C), and in a moderately coarse powder, is treated with a mixture of 90 grammes of Ether and 30 grammes of Chloroform, after brisk agitation 10 c c of a mixture of 2 parts by weight of Sodium Hydroxide Solution (15 p c) and one part by weight of Water is added, and the mixture allowed to stand for three hours, shaking well at frequent intervals. Add 10 c c, or at any rate sufficient Water to cause the powdered root to agglomerate on shaking and the supernatant liquid to clear completely. After standing for one hour 100 grammes of the clear Chloroform-Ether Solution is filtered through a dry, well-covered filter into a flask and about one-half distilled off. The remaining Chloroform-Ether Solution is introduced into a separator, the flask washed three times with a mixture of 3 parts by weight of Ether and 1 part by weight of Chloroform, and the mixed liquids well shaken with 25 c c of Centi-normal Volumetric Hydrochloric Acid Solution. After the addition of sufficient Ether to cause the Chloroform-Ether Solution to float on top of the acid liquid and after the fluids have become completely clear, the acid liquid is filtered through a small filter paper previously moistened with Water, into a flask holding about 100 c c. The Chloroform-Ether solution is shaken three times in succession with 10 c c of Water, passing the washings through the same filter paper, the latter is washed with Water and the mixed fluids diluted with Water to 100 c c. A measured quantity of 50 c c is removed, 50 c c of Water and sufficient Ether to form a layer of 1 cm. are added, and after the addition of 5 drops of Iodococci Solution, Centi-normal Volumetric Potassium Hydroxide Solution is added until the lower aqueous layer assumes a pale red coloration, shaking vigorously after each addition. Not more than 8.5 c c of

Centi-normal Potassium Hydroxide Solution should be required to effect this.

Preparations.

LINIMENTUM ACONITI. LINIMENT OF ACONITE

Powdered Aconite Root percolated with Alcohol (90 p.c.) to produce a liquid, of which 30 represents 20 of root and contains 1 of Camphor (1 in 1½)

This liniment was introduced by Peter Squire, who made it 1 in 1, and it was kept this strength in *B.P.* '64 and '67. It was diluted to 1 in 1½ in *B.P.* '85, but more recent experiments (*P.J.* '03, i 158) show that it can be made 1 in 1 and practically contain all the alkaloid.

Applied with a camel's hair pencil, alone, or mixed in equal proportions with Soap Liniment or Ammoniated Camphor Liment, and rubbed on the part (but not upon an abraded surface), relieves acute neuralgia.

TINCTURA ACONITI. TINCTURE OF ACONITE

Fr., TINCTURE D'ACONITE; Ger., ACONIT TINKTUR; Ital., TINTURA DI ACONITO; Span., TINTURA ALCOHOLICA DE ACONITO

1 of Aconite Root, in No. 40 powder, percolated with Alcohol (70 p.c.) to yield 20. (1 in 20)

Dose. 5 to 15 minims = 0.3 to 0.9 c.c.; if very frequently repeated, 2 to 5 minims = 0.1 to 0.3 c.c.

Ph. Ger. maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes; of the 1 in 10 Tincture

Tests.—Tincture of Aconite has a specific gravity of about 0.890, it contains about 1.0 p.c. w/v of total solids and about 70 p.c. w/v of Absolute Alcohol

Dr. Fleming's Tincture of Aconite (sometimes known as *Tinctura Aconiti Fortior*) was much stronger, being about the same strength as the present Liniment, 1 in 1½, but without the Camphor. **Dr. Turnbull's Tincture of Aconite** was rather weaker than Fleming's.

Foreign Pharmacopœias—Official in Mex. and Hung., 1 Root and 5 Fr., Alcoholure, 1 fresh Leaves and 1, also 1 Root in 10. Ger. and Jap., 1 Root and 10. Ital., 1 Root and 5. Port., 1 dried Leaves and 5, also 1 Root and 5, and 1 fresh Leaves and 1. Span., 1 Root in 10. All by weight. U.S., Root 10, Alcohol to measure 100. U.S. has also Fluidextractum Aconiti, 1 in 1, standardised to contain 0.4 p.c. w/v of Aconitine.

Dutch, Root percolated with Alcohol (70 p.c.) to produce a tincture containing 0.025 p.c. of alkaloids.

The *Brussels Conference* adopted a standard of 0.05 p.c. of total alkaloids, the tincture being prepared with Alcohol (70 p.c.). Belg., Fr. and Swiss adopt the *Brussels Conference* standard.

Not Official.

EXTRACTUM ACONITI RADICIS ALCOHOLICUM.—Aconite Root in powder, percolated with Alcohol (90 p.c.), and the product evaporated to a pilular consistence.

Dose.—¼ to ½ grain = 0.01 to 0.03 gramme

This must not be confounded with *Extractum Aconiti, B.P.* '85, which was made from the herb and was much weaker.

Foreign Pharmacopœias.—Official in Fr., Hung. and Russ., use 70 p.c. Alcohol; Mex., 60 p.c. Alcohol; Ital., standardised to contain not less than 0.5 p.c. of alkaloid.

CHLOROFORMUM ACONITI —Powdered Root, 20, Chloroform, to percolate, 20 Painted on with a camel's hair brush, relieves neuralgia in almost every form The above formula was that introduced by Peter Squire about fifty years ago It is preferably prepared by a similar method to that given under Chloroform Belladonna, by mixing the root in No 40 powder with Slaked Lime and powdered Ammonium Carbonate and percolating with sufficient Chloroform to produce a 1 in 1 percolate The *B P C* process employs Ammonia Solution, necessitating the use of Absolute Alcohol to allow the Chloroform to exert its full solvent action Squire's original process yielded a product having a specific gravity of 1.479, and contained 1.09 p c w/v of total solids When assayed according to the process recommended by the *U S P* for the assay of Aconite, it yielded gravimetrically 0.09 p c w/v of Aconite alkaloids yielding the same figure on titration, when assayed according to the process recommended by Farr and Wright it yielded a similar figure both gravimetrically and volumetrically Samples of the preparation prepared by the modified Squire process and by the process of the *B P C* (which consists in moistening 100 of the root in No 60 powder with 25 of Ammonia Solution, and percolating with a sufficient quantity of a mixture consisting of 1 of Absolute Alcohol to 7 of Chloroform to produce a 1 in 1 product) to some extent confirmed the results published by Farr and Wright (*P J '07*, ii 107), and showed that the former yielded a lower percentage of Aconite alkaloids than the latter A better idea of the relative efficiency of the two processes is obtained by employing the method adopted by the 8th Decennial Revision of the *U S P* for the assay of Aconite When assayed by the *U S P* process, a Chloroform of Aconite prepared by the modified Squire process showed on gravimetric determination 0.16 p c of Aconite alkaloids, which gave 0.163 p c on titration The *B P C* product assayed by the *U S P* process showed 0.264 p c by gravimetric determination and 0.266 p c on titration, when assayed by Farr and Wright's process Chloroform of Aconite prepared by the modified Squire process showed gravimetrically 0.183 p c and 0.153 p c by titration, Chloroform of Aconite prepared by the *B P C* process showed 0.26 p c gravimetrically and 0.268 p c by titration Chloroform of Aconite by the modified Squire process had a specific gravity of 1.472 and contained 0.73 p c w/v of total solids, Chloroform of Aconite prepared by the *B P C* process had a specific gravity of 1.412 and contained 1.67 p c of total solids

LINIMENTUM ACONITI COMPOSITUM (*Squire*) —Chloroform of Aconite, 1, Liniment of Aconite, 7 Sprinkled on impermeable Pline and applied for neuralgia

Linimentum Aconiti et Chloroformi —Chloroform, 12.50, Liniment of Aconite, q s to produce 100 —*B P C*.

LINIMENTUM ACONITI COMPOSITUM *Syn* A B C Liniment —Aconite Liniment, Belladonna Liniment, Chloroform Liniment, equal parts —*Guy's*

Liniment of Chloroform contains Olive Oil, which will not dissolve in the other Liniments, but it is useful as a lubricant for rubbing Some prescribers prefer to use Chloroform in place of Chloroform Liniment as follows Aconite Liniment, 2, Belladonna Liniment, 2, Chloroform, 1.

This form has been included in *B P C*.

PIGMENTUM IODI CUM ACONITO —*See* Iodine

TROCHISCI ACONITI —Each lozenge contains $\frac{1}{2}$ minim Tincture of Aconite

Dose —One Lozenge every half hour or hour in tonsillitis and febrile affections of the throat

Pastilles of Aconite are made of two strengths, $\frac{1}{2}$ minim of Tincture of Aconite and 1 minim of the Tincture Prescribers should indicate the strength required.

Emplast. Aconiti, also **Emp. Aconiti et Belladonnæ**, are made in rubber combination

ACONITINA.

ACONITINE

$C_{34}H_{47}NO_{11}$, eq 642.53

A crystalline Alkaloid, obtained from Aconite Root. It forms colourless or white odourless hexagonal rhombic prisms, which should be preserved in well-stoppered glass bottles of a dark amber tint.

Solubility.—Almost insoluble in Water, 1 in 35 of Alcohol (90 p.c.); 1 in 45 of Ether, 1 in 1 of Chloroform, soluble in Oleic Acid.

Medicinal Properties.—It relieves acute nervous pain when rubbed on the part in the form of **ointment**, producing a tingling sensation followed by numbness. Care must be taken that it does not come in contact with a mucous surface, such as the conjunctiva, or with abraded skin.

It has been applied with marked benefit in trigeminal neuralgia, and to relieve the pain of *acute* rheumatism and gout.

Dose.—As a pure crystalline Aconitine would probably be fatal to an adult in a dose of 3 milligrammes ($\frac{1}{2}$ grain), the maximum dose should not exceed $\frac{1}{10}$ milligramme ($\frac{1}{666}$ grain) *pro dosi*, or $\frac{1}{10}$ milligramme ($\frac{1}{10}$ grain) per diem, and the commencing dose should be smaller.

Solutions of the alkaloid are prone to decomposition, aqueous or alcoholic solutions should therefore be slightly acidified with **Hydrochloric Acid**, or crystallised **Aconitine Nitrate** (official in *Fr.*) should be used.

Two new Aconitines, extracted from Aconite roots indigenous to India, have formed the subject of lengthy papers in the *J. C. S. Trans.* '05, 1621, 1636. Indaconitine is the alkaloid of *Aconitum chasmanthum*, Stapf, and Bikhacointine, the alkaloid of *Aconitum speciosum*, Stapf. The pharmacology of these two alkaloids has been studied, and the conclusions arrived at (*L.* '05, n 1347) that Indaconitine and Bikhacointine may be substituted for Aconitine and Pseudaconitine for internal use. Indaconitine is admissible in the same dose as Aconitine from 4 *Napellus* and Bikhacointine in proportion of 0.75 of the unit dose of Aconitine, whilst for local application they may be used as constituents of ointments in similar proportions to the Aconitine of 1 *Napellus*.

Official Preparation.—Unguentum Aconitinum.

Not Official.—Oleatum Aconitinum.

Foreign Pharmacopœias.—Official in *Fr.*, m p 184° C.; *Ital.*, m p 186° C.; *Mex.*, m p not given; *Span.*, m p 183°; *U.S.*, m p. see below, all are crystalline products, *Hung.* specifies 'German Aconitine,' a powder, m p. 85°; *Port.*, a powder, m p. 85°. Not in the others.

Pseudaconitine.—A highly toxic crystalline alkaloid obtained from *Aconitum ferox*; only slightly soluble in Water, but readily in Alcohol and Chloroform, less readily in Ether. Dunstan gives m.p. as 201° C.—*J.C.S. Trans.* '97, 858.

Tests.—Aconitine melts at 188.5° C. (371.3° F.), 189° to 190° C. (372.2° to 374° F), *B.P.* It is distinguished by the characteristic sensation of tingling and numbness which is produced on moist mucous surfaces by extremely dilute solutions of the alkaloid or its

salts The solution of the alkaloid in Alcohol (90 p c) is dextrogyrate, whilst solutions of the salts are lævogyrate. A dilute aqueous solution of the alkaloid rendered faintly acid by the addition of Acetic Acid yields a characteristic red crystalline precipitate when Potassium Permanganate Solution is added in slight excess. On saponification Aconitine yields Acetic and Benzoic Acids and Aconine. The preparation of the Aurichloride and a determination of its melting point have been recommended as a means of identifying the alkaloid, but *BP* makes no reference to these characteristics. The melting point of pure Aconitine Aurichloride is 135.5°C (275.9°F). Aconitine is distinguished from Pseudoaconitine, Veratrine and Atropine by not yielding any violet-red coloration when the residue left on the evaporation of a small quantity with a few drops of Nitric Acid is moistened with one or two drops of an alcoholic solution of Potassium Hydroxide.

Melting Point— 388°F (195°C) on rapidly heating, *USP*, when slowly heated it decomposes and melts at 182°C (359.6°F), and on ignition it leaves no residue, *USP*.

Sulphuric or Nitric Acid—No colour is produced when Aconitine is dropped upon Sulphuric Acid or Nitric Acid, but an orange colour is produced when it is rubbed with Sulphuric Acid containing a crystal of Ammonium Vanadate, *USP*.

Potassium Permanganate Solution—Dilute solutions (up to 1-4000) of the alkaloid, when faintly acidulated with Acetic Acid, give a red crystalline precipitate with a few drops of 1 p c w/v Potassium Permanganate Solution, *BP*, the *USP* uses a 1-1000 dilution with 1 drop of Tenth normal Potassium Permanganate Solution and does not acidulate. Aconitine containing decomposition products (Amorphous Aconitine) produces this precipitate only in solutions of not less than 1-200, while Cocaine, Hydrastine and Papaverine yield similar precipitates, but only when in more concentrated solutions, *USP*.

Mercuric Potassium Iodide T.S., Tannic Acid T.S., and Gold Chloride T.S. give precipitates with dilute solutions of Aconitine, but Platonic Chloride T.S., Mercuric Chloride T.S., and Picric Acid T.S. only in concentrated solutions, *USP*.

Preparation

UNGVENTUM ACONITINÆ ACONITINE OINTMENT

Dissolve 10 grains of Aconitine in 80 grains of Oleic Acid by the aid of gentle heat, and mix with 410 grains of Lard (1 in 50).

Not Official

OLEATUM ACONITINÆ.—Aconitine, 2 grains, Oleic Acid, 98 grains, dissolve.

Dr Shoemaker states that this has a slight local action, and it can be used in mild cases of neuralgia.—*BMJ* '84, ii 750.

This has been incorporated in the *BP C* under the title **Oleinatium Aconitinæ**, *syn* Oleate of Aconitine.

ACTÆA RACEMOSA.

See CIMICIFUGÆ RHIZOMA.

ADEPS.

LARD.

FR, ANGEF, GER, SCHWEINSCHMALZ, ITAL, GRASSO SUINO, SPAN, GRASA DE CERDO

A soft white solid having a characteristic though not rancid odour and a bland taste. It is the purified abdominal fat of the Hog, *Sus scrofa*.

Adeps Induratus is Lard deprived of its oil by pressure, and the oil is known as **Lard Oil**.

Solubility.—1 in 22 of Ether, and 1 in 16 of Oil of Turpentine; almost insoluble in Alcohol (90 p.c.)

Medicinal Properties. Emollient. Added to poultices to prevent them drying and sticking to the skin.

Official Preparation. Adeps Benzoatus. Used in the preparation of Emplastrum Cichorii, Elixir Phosphoricum and the following Ointments: Acornutina, Atropine, Cocaine, Iodine, Mercury, Mercurio Sulfurico, Rosaceum and Veratrine.

Foreign Pharmacopœias. Official in Austr. (Auringia Poter.), Belg., Dan., Dutch, Fr., Ger., Hun., Ital., Jap. (Adeps, Surtinus), New (Auringia), Mex. (Manteca de Cerdo), Port. (Banha), Russ., Span., Swed., Swiss and U.S.

Tests.—The distinguishing tests for Lard are its physical appearance, melting point, specific gravity, the percentage of Potassium Hydroxide absorbed in saponification, and its Iodine absorption. *B.P.* and *P.G.* include the melting point, but no specific gravity. Melting point and specific gravity are both included in *U.S.P.* Lard melts at from 35 to 45° C. (95 to 113° F.), the *B.P.* states that it fuses at 37.8° C. (100° F.) yielding a clear liquid at a somewhat higher temperature; the *U.S.P.* gives 38 to 40° C. (100.4 to 104° F.), the *P.G.* 36 to 42° C. (96.8 to 107.6° F.). It has a specific gravity at 99° C. (210.2° F.) compared with Water at 15.5° C. (60° F.) of about 0.860, the *U.S.P.* states about 0.917 at 25° C. (77° F.) and about 0.904 at 40° C. (104° F.) compared with Water at 25° C. (77° F.). It has a Saponification value of 192 to 196.5. It has an Iodine value of 52 to 62, and should contain but a trace of free acid. Samples examined in the author's laboratory possessed a Saponification value of 192 to 197, and an Iodine value of 50 to 56. Neither *B.P.*, *U.S.P.* nor *P.G.* include the percentage of Potassium Hydroxide absorbed on saponification, the *Austr.* gives a Saponification value of 194 to 197. *P.G.*, *Ital.* and *Austr.* include an Iodine absorption, *P.G.* not less than 46 p.c. nor more than 66 p.c.; *Ital.*, 55 p.c. to 60 p.c., *Austr.*, 48 p.c. to 60 p.c., the *U.S.P.* does not include this constant.

The more generally occurring impurities are Water, Salt, an undue proportion of free Acid, Cotton Seed Oil, Beef Stearin, and Starch. No official test is given for Beef Stearin, but the Lard is required to be entirely soluble in Ether. The amount of Water may be determined by heating a weighed quantity of 5 grammes of the sample in a flat-bottomed porcelain dish at a temperature of about

105° C (221° F) A rough idea of the amount of Water may be obtained by the behaviour of the sample when shaken with Carbon Bisulphide Salt, if present, is shown by the Silver Nitrate test described below Cotton Seed Oil may be detected by the test with Alcoholic Silver Nitrate Solution given in the small type below The *BP*, *USP* and *PG* methods of determining the free acid are essentially the same, and require that when a weighed quantity of 10 grammes of the Lard is dissolved in 10 c c of Chloroform and mixed with 10 c c of Alcohol of the strength required by the respective Pharmacopæias, after the addition of 2 drops of Phenolphthalein Test solution not more than 0.2 c c of Volumetric Sodium Hydroxide Solution should be required to produce a permanent red colour This indicates a limit of 0.56 p c calculated as Oleic Acid

The test given for Cotton Seed Oil in the *BP* and *PG* is the Becchi's Silver Nitrate test, and it is still included in the *USP* for determining the presence of more than about 5 p c of Cotton Seed fats The test is preferably performed on the fatty acids and not on the Lard direct Mr F J Bevan has informed the author that a test which will readily detect even 1 p c of Cotton Seed Oil consists in heating in a salt bath for about half an hour 3 c c of the Lard with 1 c c of a 1 p c solution of Sulphur in Carbon Disulphide This test slightly modified has been included in the Eighth Decennial Revision of the *USP* to prove the absence of Cotton Seed Oil and certain other fats For the detection of Beef Stearin the microscopical appearance of the crystals separating out from the ethereal solution of the Lard is generally relied upon A portion of the sample when boiled with Water, cooled, and a few drops of Iodine Solution added, should yield no blue coloration, indicating the absence of Starch

Silver Nitrate Solution.—Distilled Water boiled with Lard, filtered, and acidulated with Nitric Acid should yield no white precipitate soluble in Ammonia Water, with T.S. of Silver Nitrate, *USP*

Alcoholic Silver Nitrate Solution.—If 5 c c of melted and filtered Lard be thoroughly mixed with a solution prepared by dissolving 0.1 gramme Silver Nitrate in 10 c c Alcohol (94.9 p c) and 2 drops of Nitric Acid, and then heated for 5 minutes on a water bath and vigorously shaken, the fatty layer on separation should not have assumed a dark reddish or brown colour, nor should there be any dark colour at the junction of the two liquids, *USP* The *BP* test is essentially the same as the *USP*, except that the *BP* prepares the 5 c c of Silver Nitrate Solution by dissolving 0.05 gramme of Silver Nitrate in 5 c c of Alcohol (90 p c) The *PG* test is made with 5 grammes of melted Lard and a solution of 0.05 gramme Silver Nitrate in 2 grammes of Ether and 10 grammes Alcohol, and the mixture is warmed for 15 minutes on the water bath

Sulphur in Carbon Bisulphide.—2 c c of melted and filtered Lard are mixed with 1 c c each of Amyl Alcohol and a 1 p c solution of Sulphur in Carbon Disulphide in a test tube The test tube is immersed to one-third or half its depth in boiling salt Water No reddish colour should develop in the mixture in from 10 to 15 minutes, *USP*

Saponification.—A mixture of 2 parts Lard, 3 parts Potassium Hydroxide Solution, and 2 parts of Alcohol, when boiled until it clarifies, should produce, on the addition of 50 parts of Water and 10 parts of Alcohol, a clear or only faintly opalescent liquid, *PG*.

Preparation.**ADEPS BENZOATUS. BENZOATED LARD**

Digest 210 grains of Benzoin in 16 oz of melted Lard on a water-bath for two hours, strain, and stir whilst cooling (1 in 33)

Benzoated Lard should not be used for **eye ointments** as it is irritating

Benzoated Suet (*Sevum Benzoatum*) *Ind* and *Col Add* should be used in India, in place of Benzoated Lard

Official Preparations.—Following Ointments: *Belladonna*, *Cantharides*, *Chrysarobin*, *Iodide*, *Mercuric Oleate*, *Mercurous Chloride*, *Potassium Iodide*, *Stavesacre*, *Sulphur*, *Sulphur Iodide*, and *Zinc*.

Foreign Pharmacopœias Official in Dan, Dutch, Ital. (*Grasso con Benzoino*), Norw., Swed and U.S., Benzoin 2, Lard 100, Russ., Benzoin 2, Lard 100, dried Sodium Sulphate 5, Mex. (*Pomada Benzoadæ*), Tincture of Benzoin 5, Lard 100, Fr., Benzoin 3, Lard 100, Ger. and Jap., 1 Acid Benz. in 100; Swiss, Lard 100, Benzoin 1, Dried Sodium Sulphate 6; or Lard 10, Ethereal Tincture of Benzoin 1. Not in the others.

Not Official.

UNNA'S SALVE MULLS. The bases of these are hog's lard and beef suet (singly or combined), with which are incorporated various medicaments, spread on muslin

ADEPS LANÆ.**WOOL FAT**

Fr., *GRAISS DE LAINE*, Ger., *WOLLEFETT*, Ital., *LANOLINA*, Span., *LANOLINA*

A pale yellow, tenacious, unctuous product possessing a faint characteristic odour. It is the purified fat of sheep's wool, consisting chiefly of Cholesterol, Iso cholesterol, and their Esters.

Solubility Readily soluble in Chloroform and Ether, but only partially so in Alcohol (90 p.c.) Its own weight of Water can be incorporated with it.

Medicinal Properties.—Emollient, is very readily absorbed by the skin, and thus promotes the action of remedies combined with it.

Official Preparation *Adeps Lanæ Hydrostus*.

Not Official.—*Unguentum Lanolin*, *Unguentum Adipis Lanæ*, *Adeps Lanæ cum Oleo*.

Foreign Pharmacopœias Official in Austr., Dan., Dutch, Fr., Ger., Ital. (*Lanolina*), Jap., Russ., Span., Swed., Swiss and U.S. Not in the others.

Tests.—The distinguishing tests for *Adeps Lanæ* are the melting point, which should be about 40° C. (104° F.), and the formation of the purple-red colour of Cholesterol when its solution in Chloroform is poured gently on to the surface of Sulphuric Acid. It is preferable to use an acid containing a trace of Water for performing this test.

The *B.P.* gives the melting point as 40° to 44·4° C. (104° to 112° F.); the *U.S.P.* and the *P.G.* about 40° C. (104° F.); the *U.S.P.* states that at a higher temperature it vaporises and burns with a luminous

sooty flame, the *P G* that Wool Fat burns with a luminous very smoky flame

The percentage of Potassium Hydroxide required for saponification, as based upon Helbing's saponification test, affords a good criterion of the purity of the sample, but no official mention is made of this test. A thoroughly purified Wool Fat will combine with about 8 to 9 p c of Potassium Hydroxide, Glycerin Fats give much higher figures (Lard, 19.5 p c, Olive Oil, 18.0 p c, Cocoa Nut Fat, 26 p c), while Petroleum bases, being unsaponifiable, do not consume any.

The more generally occurring impurities are mineral matter, an undue amount of free acid, and nitrogenous organic matter. The *BP*, *USP* and *P G* all require that the residue left on ignition shall not be alkaline to Litmus, but differ in the amount of mineral residue permissible. *BP* and *USP* state not more than 0.3 p c, *P G* allows, at the highest 0.5 p c. The *BP* does not allow more than 0.28 p c of free Acid calculated as Oleic Acid, *P G* not more than 0.14 p c, and *US* not more than 0.7 p c. In carrying out the determination of the free acid, the *BP* uses 25 c c Ether as a solvent for the 10 grammes of Wool Fat and employs Volumetric Sodium Hydroxide Solution, of which not more than 0.1 c c should be required, the *USP* and *P G* use 2 grammes of the Wool Fat for the determination, employing 10 c c of Ether as a solvent, the *USP* uses Normal Volumetric Potassium Hydroxide Solution for the titration and the *P G* Tenth-normal Volumetric Potassium Hydroxide Solution, the *USP* stipulates that only 1 drop of the Normal Volumetric Solution shall be required, the *P G* 0.1 c c of Tenth normal Volumetric Potassium Hydroxide. The *BP* does not include a test for the absence of Glycerin, but such a test is included in both the *USP* and *P G*, as well as a test for ensuring the absence of Chlorides, due to Chlorine substitution products formed during the bleaching process. In carrying out the test for Glycerin both the *USP* and the *P G* work upon the filtrate obtained by shaking the Wool Fat with boiling Water, both Pharmacopœias employing 10 grammes of the Wool Fat and 50 c c of Water, and requiring that the clear separated aqueous liquid shall not yield a residue of Glycerin on evaporation, nor vapours of Ammonia when boiled with a solution of an alkali Hydroxide, the *USP* employing Potassium Hydroxide T.S., the *P G* Calcium Hydroxide Solution, the latter portion of the test is intended to detect nitrogenous organic matter. In carrying out the test the *BP* boils the Wool Fat itself with Sodium Hydroxide Solution. The *P G* tests another portion of 10 c c of the clear separated aqueous liquid with 2 drops of Potassium Permanganate Solution (0.1 p c w/w), requiring that the mixture shall maintain its red colour. This test is intended as confirmatory evidence of the absence of Glycerin and other readily oxidisable organic impurities. The absence of Chlorides is determined by the test with Alcoholic Silver Nitrate Solution given in the small type below.

Sulphuric Acid.—If a solution of Wool Fat in Chloroform (1.50 *P G* and *USP*) be poured as a layer over Sulphuric Acid, a zone of deep brownish red

colour gradually appears at the line of contact of the two liquids, *P.G.* and *U.S.P.* (the *B.P.* gives no quantities and states simply in colour), it attains its deepest shade after about 24 hours, *P.G.*

Alcoholic Silver Nitrate Solution.—1 gramme of Wool Fat boiled with 20 c.c. Alcohol, and filtered, yields a filtrate which on cooling should not be rendered turbid by alcoholic Solution of Silver Nitrate (1-20), *P.G.*, *U.S.P.*, or a turbidity disappearing on warming, *P.G.*

Preparation.

ADEPS LANÆ HYDROSUS. **HYDROUS WOOL FAT**

A nearly white, or yellowish white, unctuous mass prepared by incorporating 3 of Distilled Water with 7 of Wool Fat by rubbing together in a warm vessel. Used as a basis for ointments. It does not become rancid. Mixes with about half its weight of Water. It is better for ointments when mixed with an equal weight of Soft Paraffin.

Official Preparations. Used in the preparation of Unguentum Comi, and Unguentum Hamamelidis.

Foreign Pharmacopœias. Official in Anti (Adeps Lanae Hydrosus), Norw (Lanolinum Hydratum), Ital and Mex (Lanolina), Dan., Dutch, Ger., Jap and Russ (Adeps Lanae cum Aqua), Swed (Adeps Lanae), Swiss (Lanoline), U.S. Not in the others.

Tests.—Hydrous Wool Fat melts at about 40° C. (104° F.) separating into two layers, an upper oily layer and a lower aqueous layer. It is officially required to yield not less than 70 p.c. of residue when dried till constant in weight at a temperature of about 100° C. (212° F.), thus indicating a loss of not more than 30 p.c. This is also the *U.S.P.* limit, the latter Pharmacopœia adding in addition that the yellowish tenacious unctuous mass remaining should be completely soluble in Ether or Chloroform, and only sparingly soluble in Alcohol, should respond to the tests given under Adeps Lanæ. The *P.G.* requires that at the temperature mentioned it shall not lose more than 26 p.c. of its weight, and that the residue remaining after the separation of the Water shall answer the tests for Wool Fat and may be tested for its purity by similar methods to those employed for the anhydrous fat.

Not Official.

UNGUENTUM LANOLINI.—Hydrous Lanoline, 2, Soft Paraffin, 1

UNGUENTUM ADIPIS LANÆ (*Ger.*) -- Wool Fat (anhydrous), 20, Water, 5, Olive Oil, 5. All by weight.

An improvement on this is the following:—

ADEPS LANÆ CUM OLEO. Hydrous Wool Fat, 9, Almond Oil, 1.

This has been incorporated in the *B.P.C.* as follows:—Hydrous Wool Fat, 10, Olive Oil, 10, under the title Unguentum Adipis Lanæ.

Not Official.

ADHATODA.

The fresh and the dried leaves of *Adhatoda Vasica*, are official in *Ind* and *Col Add* for India and the Eastern Colonies as are also **Extractum Adhatodæ Liquidum** (1 in 1), dose 20 to 60 minims=1·2 to 3·6 c.c., **Succus Adhatodæ**, the freshly expressed and strained juice of the bruised fresh leaves, dose 1 to 4 fl. drms=8·6 to 14·2 c.c., and **Tinctura Adhatodæ** (1 in 8), dose 30 to 60 minims=1·8 to 3·6 c.c.

Not Official.

ADONISThe leaves and stalks of *Adonis vernalis*, L

Medicinal Properties—A cardiac tonic Useful in mitral and aortic regurgitation, relieving intracardiac pressure and præcordial pain—*L* '88, 11 1012 A useful adjunct to bromides in epilepsy—*L* '94, 11 1288, *B M J E* '95, 1 12, and '98, 1 44

Dose—2 to 6 grains=0.13 to 0.4 gramme in powder, or the equivalent of an infusion, tincture, or fluid extract

Foreign Pharmacopœias—Official in Ital, Russ, Span and Swiss Not in the others

ADONIDIN—A glucoside, very deliquescent, soluble in Water and Alcohol (90 p c)

Dose— $\frac{1}{2}$ to $\frac{1}{2}$ grain=0.01 to 0.03 gramme Generally given in pill.

ADRENALIN—*See* SUPRARENAL GLAND.

ETHER.**ETHER**

FR, ETHER, GER, AETHER, ITAL, ETHERE, SPAN, ETHER

A light, colourless, volatile, mobile liquid, possessing a strong characteristic odour and containing not less than 92 p c by volume of Ethyl Oxide ($C_2H_5)_2O$, eq 73.52 It is also known as Ethyl Ether, and Sulphuric Ether It is very volatile, and gives off a very inflammable and very heavy vapour

It is prepared by the action of Sulphuric Acid upon Alcohol, and subsequent distillation and rectification of the product

This product contains both Alcohol and Water, but *Æther Purificatus* (*see* below) is almost free from both

Solubility.—1 in 10 of Water, mixes in all proportions with Alcohol (90 p c)

B P states that it is miscible in all proportions with Chloroform, but the mixture forms a turbid liquid, owing to the presence of Water in the Ether

Water dissolves a tenth of its volume of Ether, and reciprocally Ether takes up about the same proportion of Water

Ether dissolves Corrosive Sublimate, Red Mercuric Iodide, Iodine and Bromine freely, Sulphur and Phosphorus sparingly It is also a solvent of the volatile and fixed oils, many resins and balsams, caoutchouc, and most of the organic vegetable alkaloids It does not dissolve Potassium or Sodium Hydroxides, in which respect it differs from Alcohol

Medicinal Properties.—It is a rapid and powerful diffusible stimulant, antispasmodic and carminative, and is of great use in syncope or heart-failure from any cause, dyspnoea, gastralgia, flatulence, spasmodic asthma and angina pectoris It excites secretion from the mucous surfaces of the alimentary tract, and, as it stimulates the pancreas, it is sometimes given with Cod Liver Oil

As an anæsthetic, *see* *Æther Purificatus* and *Æther Methylicatus*.

Dose.—For repeated administration, 10 to 30 minims = 0·6 to 1·8 cc; for a single administration, 40 to 60 minims = 2·4 to 3·6 cc

When used hypodermically for heart failure the dose is 15 to 30 minims = 0·9 to 1·8 cc

Prescribing Notes.—*Best prescribed as Spirit of Ether, which mixes readily with Water. 'Perles' are prepared*

Official Preparations. Ether Purificatus, Spiritus Ætheris, Spiritus Ætheris Compositus. It is the preparation of Colloidium, Extractum Filicis Jaquandii. Ether Purificatus is used in the preparation of Extractum Staphylini, and Spiritus Ætheris in Tinctura Lobeliae Ætherea

Not Official. Ether Methylatus, Mistura Ætheris cum Ammonia, Sirop d'Ether, Spiritus Ætheris Muraticus

Foreign Pharmacopœias. See under Ether Purificatus.

Tests.—The distinguishing tests for Æther are its volatility as indicated by its boiling point, the specific gravity, and the odour. It has a boiling point of about 40° C (104° F); the *BP* gives the boiling point as below 40·5° C (105° F). Æther sp. gr 0·735 has been official in Great Britain for many years, but Foreign Pharmacopœias only recognise the fluid which is official in the *BP* as Ether Purificatus, see below

The more generally occurring impurities are an undue amount of Acid, or of Alcohol, extractive matter, and organic impurities produced during its manufacture and which have escaped separation during the process of rectification

It should not possess an acid reaction towards blue Litmus paper, indicating the absence of acidity. When shaken with Water the volume should not be decreased by less than one-tenth, indicating the absence of excess of Alcohol. It should be completely volatilised without leaving a residue, indicating the absence of extractive matter; when mixed with an equal volume of Sulphuric Acid, the mixture kept well cooled, it should yield little or no coloration, indicating the absence of organic impurities

ÆTHER PURIFICATUS. PURIFIED ETHER

A colourless, transparent, very volatile, mobile and inflammable liquid from which the greater part of the Alcohol has been removed by washing with Water, and the Water by distillation over a dehydrating agent. *U.S.P.* describes it as containing about 96 p.c. by weight of Ethyl Oxide

It is the *Æther pro narcosi* of the German and Swedish Pharmacopœias and the official Æther of the *U.S.*

Medicinal Properties.—Used for producing general anæsthesia by inhalation.

It has also been used as a spray for obtaining local anæsthesia in minor surgery, and to relieve severe neuralgic pain, but as the lower the boiling point of the Ether the more complete is the anæsthesia, Methylated Ether, sp. gr. 0·717 (see below), is preferable for use with the spray.

Æther was first used as an anæsthetic for capital operations in 1846, and

Purified Ether is preferred by some to Chloroform, as it has a less depressing effect upon the heart, vessels, and respiratory centre. It is used also in conjunction with Nitrous Oxide for minor operations in dentistry and surgery.

Official Preparation—Used in the preparation of *Extractum Strophanthi*.

Foreign Pharmacopœias—Official in Austr., sp gr 0.720, Russ., sp gr 0.725, Belg., Dutch, Fr., Ger. and Jap., sp gr 0.720, Fr., also sp gr 0.724, Hung., sp gr 0.724 to 0.728, Port., sp gr 0.728, Span. (Ether), sp gr 0.720, Dan., Norw., Ital. (Ether), Swed. and Swiss, sp gr 0.720 to 0.722, U.S., sp gr 0.716 to 0.717 at 25° C (77° F), Mex. (Ether Sulfurico), sp gr 0.720.

Tests.—The distinguishing tests for Æther Purificatus are its boiling point, its specific gravity, its peculiar odour and physical appearance. The *BP* does not give a boiling point, but requires that it should not commence to distil below 34.5° C (94.1° F), the *USP* gives the boiling point at 35.5° C (96° F), the *PG* as 35° C (95° F). The specific gravity given in the *BP* is from 0.720 to 0.722, that of the *USP* 0.716 to 0.717 at 25° C (77° F), that of the *PG* 0.720.

The more generally occurring impurities are Methylc Ether, extractive matter, excess of Water, Aldehyde, acid, organic impurities, and Hydrogen Peroxide. Methylc Ether may be detected, if present, by the lowering of the boiling point. Both the *USP* and the *PG* require that Ether shall leave no residue on evaporation, indicating the absence of extractive matter. The *PG* states that when allowed to evaporate at ordinary temperature it leaves a damp ring which should not redden blue Litmus paper. The three Pharmacopœias differ considerably in their methods of testing for Aldehyde, Hydrogen Peroxide, and excess of Water. All three employ Potassium Hydroxide as a test for Aldehyde, the *BP* and *PG* use the solid form, *USP* the solution. The *BP* gives neither a time limit within which no coloration shall be produced, nor quantities, *USP* uses 10 of Ether to 1 of Potassium Hydroxide Solution and a time limit of 'within one hour,' whilst *PG* states no quantity for the test, but requires that no yellow colour shall be produced within six hours, the mixture being protected from the light.

In testing for Hydrogen Peroxide, *BP* uses Potassium Bichromate acidified with Sulphuric Acid, requiring that the ethereal liquid shall not develop a blue colour, the *PG* employs Potassium Iodide Solution and protects the mixture from the light, no coloration should be produced within one hour, whilst the *USP* does not include a test for Hydrogen Peroxide. Excess of Water is officially detected by mixing equal volumes of Ether and Carbon Bisulphide, when a clear solution should be obtained, in the *USP* the test is performed with Ether-saturated Water, and is made quantitative by employing equal volumes of the menstruum and Ether, and noting the decrease in volume of the ethereal liquid, 20 c.c. of the sample shaken with 20 c.c. of Ether-saturated Water should not measure less than 19.2 c.c. The *PG* does not include a test for excess of Water.

Odorous impurities, if present, may be detected by the odour imparted to clean, odourless filter paper when the Ether is allowed to spontaneously evaporate. Neither the *BP* nor the *PG* states the quantity to be used for the test, the *USP* evaporates 10 c.c. in

portions, all three Pharmacopœias require that the filter paper, after the evaporation of the Ether, shall possess no odour.

Evaporation Clean, odourless filter paper moistened with Ether, and the Ether allowed to spontaneously evaporate, should be free from any foreign odour when the Ether has evaporated, *B P*, the *P G.* and *U S P* allow 10 c.c. to evaporate.

Preparations.

SPIRITUS ÆTHERIS. SPIRIT OF ETHER The *HOFFMANN'S* ANODYNE of the Continental Pharmacopœias.

Ether, 1, Alcohol (90 p.c.), 2 (1 in 3)

Specific Gravity.—0.806 to 0.811.

Dose. For repeated administration, 20 to 40 minims = 1.2 to 2.4 c.c.; for single administration, 60 to 90 minims = 3.6 to 5.4 c.c.

Foreign Pharmacopœias Official in Austr., Dan., Ger., Hung., Jap., Norw., Swed. and Switz., 1 and 3, Belg. (Æther Sulphuricus Alcoholicus), 468 in 1000, Dutch (Æther cum Spiritu), 1 and 1, sp. gr. 0.777 to 0.782, Fr. (Æther Alcoholici), 1 and 1, sp. gr. 0.783; Ital. (Liquore Anodino di Hoffmann), 1 and 1, Mex. (Licor de Hoffmann), Æther 1, Alcohol 90 p.c., 1, Port. (Æther Alcoholizado), 7 and 3; Russ., 1 and 2, sp. gr. 0.800, Span. (Eter Sulfurico Alcoholizado), 4 and 1; U.S., $3\frac{1}{2}$ in 10. All by weight, except U.S.

Dan., Norw. and Swed. include also Æther Spirituosus Camphoratus, containing 15 p.c. of Camphor.

SPIRITUS ÆTHERIS COMPOSITUS. COMPOUND SPIRIT OF ETHER *B P Syn* - *HOFFMANN'S ANODYNE*

Ether, $5\frac{1}{2}$; Alcohol (90 p.c.), 38; and an undefined quantity of ethereal liquid, obtained by the action of 36 of Sulphuric Acid on 40 of Alcohol (90 p.c.), and subsequent treatment.

The official directions are founded on the formula of the old Dublin Pharmacopœia.

Dose For repeated administration, 20 to 40 minims = 1.2 to 2.4 c.c., for a single administration, 60 to 90 minims = 3.6 to 5.4 c.c.

Foreign Pharmacopœias Official in U.S., Ether, 325, Alcohol, 650; Ethereal Oil, 25. Not in the others.

Tests - The distinguishing tests are the sp. gr. which should be about 0.810, and the production of an opalescent mixture on the addition of Water. The more generally occurring impurities are those of an empyreumatic nature, having their origin in the process of manufacture, and which may be detected by any objectionable odour imparted to the residue left on the spontaneous evaporation of a few c.c. of the sample.

Not Official.

ETHER FROM METHYLATED SPIRIT. *Syn* Æther Methy-
latus, METHYLATED ETHER—Sp. gr. 0.717. It can be purified to such an extent by washing and redistillation as to be scarcely distinguishable from that made from pure Spirit. The Methylæ Ether being so extremely volatile is almost wholly lost during the purification.

An Ether, sp gr 0.715, can be obtained in limited quantity by careful working, occasionally samples are drawn over at 0.713, in cold weather

Medicinal Properties—It is largely employed as a **spray** for local anæsthesia, as well as for **inhalation**. As in the case of 'Methylated Chloroform,' the impurities from the Wood Spirit, employed in the manufacture, can be completely eliminated

Ether can be made more volatile for use with the **spray** by the addition of 20 per cent of a light Petroleum Ether

Ether from Methylated Spirit, sp gr 0.720 washed and redistilled, is also supplied for **inhalation**. It is *not* so suitable as the above for the **spray** because it volatilises less rapidly

MISTURA ÆTHERIS CUM AMMONIA—Spirit of Ether, 30 minims, Aromatic Spirit of Ammonia, 30 minims, Distilled Water, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

Mistura Ammoniae et Ætheris—Aromatic Spirit of Ammonia, 30 minims, Spirit of Ether, 30 minims, Chloroform Water, to make 1 oz—*St Mary's*

SIROP D'ETHER SIROPUS CUM ÆTHERE—Ether (sp gr 0.720), 2, Distilled Water, 28, Alcohol (90 p c), 5, Syrup, 70, all by weight—*Fr*

This has been incorporated in the *B P C* under the title **Syrupus Ætheris Compositus**

SPIRITUS ÆTHERIS MURIATICUS *Syn*—*SP* **SALIS DULCIS**, CLUTTON'S FERRIFUGÆ SPIRIT

A colourless liquid Sp gr 0.860

A very old preparation, still prescribed for pyrexia, and cold in the head

Dose—30 to 60 minims = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Norw (*Æther Chloratus Spirituosus*)

ÆTHER ACETICUS.

ACETIC ETHER

FR, ACÉTATE D'ETHYLE, *GER*, ESSIGÄTHER, *ITAL*, EIFFRE ACETICO, *SPAN*, ÉTER ACÉTICO

A transparent, colourless, volatile, inflammable liquid, possessing a characteristic ethereal odour and taste. It consists almost entirely of Ethyl Acetate, $C_2H_5C_2H_3O_2$, eq 87.40, with a small quantity of Ethylic Alcohol, Water and possibly traces of organic impurities

A good commercial specimen should contain 90 p c of Ethyl Acetate and about 10 p c of Ethylic Alcohol

It should be kept in well stoppered, amber-tinted bottles, and in a cool atmosphere

Solubility.—About 1 in 9 of Water. Pure Acetic Ether is miscible in all proportions with Alcohol (90 p c), and with Ether, but Acetic Ether of *BP* specific gravity is not soluble in all proportions in Chloroform, but if such Ether be dehydrated over Calcium Chloride or Potassium Carbonate it will then mix with Chloroform in all proportions, but the purified product has a specific gravity of 0.895.

Medicinal Properties. — Antispasmodic, stimulant, and carminative. It is also used as a sedative **inhalation** in irritation of the larynx, 30 minims in a pint of Water.

Dose. For repeated administration, 20 to 40 minims = 1 2 to 2 4 c.c., for a single administration, 60 to 90 minims = 3 6 to 5 4 c.c.

Official Preparation. — Used in the preparation of Liqueur Epispasticus

Foreign Pharmacopœias Official in Austr., Belg., Dan., Dutch, Ger., Jap., Norw., and Russ., sp. gr. 0·900 to 0·904, Hung., sp. gr. 0·900; Belg., sp. gr. 0·890. Fr., sp. gr. 0·920. Ital. (Ethere Aceticæ), sp. gr. 0·900 to 0·904, Mex. (Ether Aceticæ) sp. gr. 0·920; Port., sp. gr. 0·920, Span., sp. gr. 0·915, Dan. and Swed., sp. gr. 0·902 to 0·906, Swiss, sp. gr. 0·904; U.S., sp. gr. 0·883 to 0·885 at 25° C. (77° F.)

Tests The distinguishing tests for Acetic Ether are its sp. gr. of about 0·900, its boiling point, which should be about 162° C. (323·6° F.), and the limits of temperature between which it distils, 165° and 172° F. (73·9° and 77·8° C.)

It is officially required to have a specific gravity of 0·900 to 0·905; the *U.S.P.* gives 0·883 to 0·885 at 25° C. (77° F.), the *P.G.* 0·900 to 0·904. The official Acetic Ether is the fraction boiling between 73·9° and 77·8° C. (165° and 172° F.), the *U.S.P.* gives the boiling point of Acetic Ether from 72° to 77° C. (161·6° to 170·6° F.), the *P.G.* 74° to 76° C. (165·2° to 168·8° F.).

The more generally occurring impurities are Ethylic Alcohol, Water, free Acetic Acid, readily carbonisable organic impurities, and Ethers other than Acetic derived from impurities in the Ethylic Alcohol used in the process of manufacture. Except by inference the *B.P.* omits a test for excess of Ethylic Alcohol and Water, but both *U.S.* and *P.G.* stipulate that it shall not contain more than 10 p.c., as ascertained by the decrease in volume of the ethereal layer or the increase in volume of the aqueous layer, when a saturated aqueous solution of Acetic Ether is shaken with an equal volume of the specimen under examination. A free acid test performed either with Litmus Solution or paper is common to all three Pharmacopœias, as is also a test with Sulphuric Acid for readily carbonisable organic impurities. The latter test is included in all three Pharmacopœias, but in performing it the *B.P.* simply directs the liquids to be mixed, the *U.S.P.* and the *P.G.* direct that the Acetic Ether be carefully poured as a layer on the acid, and no coloured zone should be produced at the point of contact.

The *B.P.* and *P.G.* require that when evaporated from filter paper no odour shall remain, but do not mention the specific impurities indicated by the test. *U.S.*, on the other hand, state that the final odour should not resemble that of propyl ether, meaning the absence of Butylic and Amylic Ethers. It should be completely volatile, and should leave no weighable residue on evaporation.

ÆTHERIS NITROSI SPIRITUS.

See SPIRITUS ÆTHERIS NITROSI

Not Official.

ÆTHYL BROMIDUM

ETHYL BROMIDE HYDROBROMIC ETHER

FR, BROMURE D'ETHYLE, GER, ÄTHYLBROMID, ITAL, BROMURO DI ETILF,
SPAN, ÉTER BROMHÍDRICO C_2H_5Br , eq 108 17

A heavy, colourless, mobile and very volatile liquid, which should be preserved in well stoppered dark amber tinted bottles. It is best prepared by acting upon Potassium Bromide with Sulphuric Acid in the presence of Alcohol. Its liability to decomposition may be prevented by exclusion of light and air, and by the addition of Alcohol, which lowers the sp gr.

Shown that the process recommended in the *Companion* (17th edit) is the correct one. 60 p.c. of the commercial samples stated to be quite unfit for use — *P J* '02, i 491

Solubility — 1 in 120 of Water, but will vary with sp gr of sample, it mixes in all proportions with Alcohol (90 p.c.) and Ether.

Medicinal Properties — It is a local and general anæsthetic, more rapid in its action than Chloroform, and occasionally used in conjunction with it. It is useful in minor surgery, also in obstetric practice and in dental operations.

It should be administered in the same manner as Ether, it is very prompt in its action. It should not be given in prolonged operations or in renal disease. Has been used as a spray to produce local anæsthesia.

Strongly recommended in dental operations.

Recommended as a general anæsthetic in short operations. Action rapid, and particularly well adapted for children. Dangerous when administered with air, or if administration is protracted. Amount required varies up to 3 fl drms = 10 6 c.c. — *L* '99, ii 850, *B M J* '02, ii 589

Administered by inhalation in a single dose of from 15 to 80 grammes was free from risk, but prolonged administration of repeated doses was dangerous. More than one warning, however, has been given that 80 grammes (6 fl drms) is too large and may occasionally give rise to unpleasant symptoms — *L* '03, ii 745

A solution, 1 in 200 of Water, in angina pectoris, dose $\frac{1}{2}$ to 2 oz = 14 2 to 56 8 c.c. — *M A* '87, 24

Foreign Pharmacopœias — Fr (Bromure d'Ethyle), Dutch and Swed (Brometum Aethylicum), Belg, Ger and Swiss (Äther Bromatus), Ital (Bromuro di etile), Mex and Span (Éter Bromhídrico), Jap and Russ (Æthylum Bromatum). Not in the others.

Tests — Pure Ethyl Bromide has sp gr 1 473. Sp gr given in *P G* is 1 458 to 1 457. A very pure sample sold as containing 1 p.c. of Alcohol had sp gr 1 461, but ordinary samples may run as low as 1 94.

The boiling point of pure Ethyl Bromide is 101 8° F (38 5° C)

Boiling point of a sample sp gr 1 45 was 38 5° C (101° F), and dissolved in 120 parts of Water.

It should give no reaction with pure Sulphuric Acid, or no more than a yellow colour after an hour, indicating the absence of organic Sulphur compounds, Amyl and Ethylene compounds. When evaporated should leave no residue, indicating the absence of fixed residue. When shaken with an equal volume of Water it should be a little, if anything, decreased in volume, indicating the absence of more than a trace of Alcohol, and when the aqueous layer is separated it should not be acid in reaction towards Litmus paper, nor should an immediate turbidity be produced on the addition of a few drops of Silver Nitrate Solution, indicating the absence of Hydrobromic Acid, etc. When 1 c.c. of Ethyl Bromide is warmed with 8 drops of Aniline and 2 c.c. of Alcoholic Potassium Hydroxide Solution the characteristic odour of Carbylamine should not be evolved, indicating the absence of Chloroform. Its vapour should have a pleasant ethereal odour.

ÆTHYLENE BROMIDE, $C_2H_4Br_2$ — A heavy, colourless, somewhat volatile liquid, obtained from the hydrocarbon Ethylene. Boils at 264 2° F. (129° C.), sp gr 2 17. Given in epilepsy.

Dose — 1 to 2 minims = 0 06 to 0 18 c.c., dissolved in oil.

Not Official.

ÆTHYL CHLORIDUM.

ETHYL CHLORIDE HYDROCHLORIC OR MURIATIC ETHER

Fr, CHLORURE D'ETHYLE, GER., CHLORÄTHYL, ITAL, CLORURO D'ETILE



A colourless ethereal, inflammable liquid, which is supplied in glass capsules closed by a screw or spring cap.

It has a characteristic ethereal and somewhat agreeable odour and a sweetish burning taste.

It is produced by the action of Hydrochloric Acid gas on absolute Ethylic Alcohol.

On account of its highly inflammable and volatile nature it should be preserved in hermetically sealed glass tubes, and should not be opened when near a naked flame.

Medicinal Properties. Used for producing local anaesthesia in minor surgery and dentistry and as an analgesic in neuralgic and rheumatic pains - - *T* '03, '93, '87, '91, 119. See also Methyl Chloride.

Description of an apparatus for administering volatile anæsthetic agents, such as bromide and chloride of ethyl. A mask which hermetically seals the mouth and nose of patient - *L* '02, n 19. Caution required in the use thereof, such volatile bodies are depressants, and their capacity for danger culminates when mouth and nose are hermetically sealed - *L* '02, n 171.

A few cases of ethyl chloride narcosis. Its use recommended in place of nitrous oxide gas - *L* '01, n 699, '01, n 123. Treatment of lupus with ethyl chloride - *B M J* '01, n 76, *T G* '01, 603.

Review of 150 cases in which it was used as a general anæsthetic, with no trouble during narcosis ascribable to the drug itself - *L* '03, n 952.

A record of 100 successful administrations each of Ethyl Chloride and of Scornnoform alone, and in mixture with Nitrous Oxide. The dose was generally determined by the amount of operative work necessary in each individual case, the largest dose given being 6 c.c. - *L* '05, n 1176. A dose of 5 c.c. may be trusted (*B M J* '05, n. 616) to give at least 5 minutes' anæsthesia in midwifery cases, and gives great command over uncontrollable pains at the end of the second stage.

As an anæsthetic for young children aged from 5 days old and upwards, it is regarded (*L* '05, n 1542 1922) as one of the best means of procuring an anæsthesia of from 5 to 15 minutes. A celluloid inhaler is used, and the dose employed for infants of a few days or a few weeks old is 3 c.c., for those of 6 months and upwards 5 c.c.

An examination recorded (*L* '05, n 1691) of seven makes of this liquid obtainable in London showed that all the branded samples were pure, as was also one of the unbranded samples. One branded and one unbranded sample were found to contain traces of impurity, but the former was not intended for general anæsthesia. The boiling point of the pure product is 12.5° C (54.5° F.). It should be free from Water from foreign chlorides, from acids, from aldehydes, from Ether or Alcohol, and from organic substances.

The death of a patient (*L* '06, n 544 *B M J* '06, n. 544) while under its influence for the purpose of having four teeth removed gave rise to considerable discussion regarding its safety. The statistics published (*B M J* '06, n 616) indicate that it is not so innocuous as previous observations appeared to indicate. The dangers seem formidable (*B M J* '06, n 1007) but should not blot out its undeniable advantages. It occupies a distinct place in the armamentarium of the anæsthetist, it conveniently and safely leads up to Ether narcosis and may take the place of Nitrous Oxide when this is not available, not efficacious or unsatisfactory. The dose recommended is 2 or 3 c.c. for a child, 3 or 4 c.c. for a woman or weakly man, and 4 or 5 for a man. Minimal dosage and not too great deprivation of air when anæsthesia is complete are the patient's safeguards.

It is stated (*L* '05, n 1026) to occupy a position as an anæsthetic midway between Nitrous Oxide and Ether. It is an ideal preliminary to etherisation,

and more especially for great smokers, alcoholics, and nervous patients. Patients with obstructed breathing are regarded as bad subjects for its administration. Mixtures of Ethyl Chloride, Ethyl Bromide, etc., appear to possess no advantages.

Although in point of safety (*L* '05, ii 1922), Ethyl Chloride, when administered by skilled hands, as a general anæsthetic, comes between Ether and Chloroform, it is far from being an absolutely innocuous anæsthetic.

A record of twenty two fatalities which have occurred under Ethyl Chloride — *L* '06, i 1233.

A remarkable uniformity (*L* '06, ii 106) is noticed in the qualitative effects of Ethyl Bromide, Chloride, and Iodide. Their physiological action differs in degree only, depending simply on the volatility of the drugs. Their action on the circulatory system appears to be almost directly upon the heart.

Acid intoxication following Ethyl Chloride anæsthesia — *L* '04, i 284.

Tests — Sp gr 0.921 at 0° C (32° F). It is but slightly soluble in Water, but mixes in all proportions with Alcohol (90 p.c.), Chloroform, and Ether. It boils at 12° C (53.6° F). On ignition it burns with a green edged, smoky flame producing Hydrochloric Acid gas. It should volatilise completely at ordinary temperatures without leaving a residue, and if allowed to evaporate from pure filter paper should leave no unpleasant odour. Its aqueous or alcoholic solution should have no acid reaction towards blue Litmus paper, nor should a turbidity be produced by the addition of Silver Nitrate Solution.

U.S.P. requires that no odour of Aldehyde shall be developed when the aqueous layer, which remains after shaking 10 c.c. of Ethyl Chloride with 10 c.c. of cold Water and spontaneously evaporating the Ethyl Chloride layer, is treated with Potassium Dichromate and diluted with Sulphuric Acid, indicating the absence of Alcohol.

Foreign Pharmacopœias — Austr (Aethylum Chloratum), Belg and Swiss (Æther Chloratus), Dutch (Chlorietum Aethylicum), Fr (Chlorure d'Éthyle), Ital (Cloruro d'Etil), U.S.

NARCOTILE (Methylene bichloride). A transparent, colourless, mobile, highly volatile and inflammable liquid. Introduced as a new general anæsthetic — *L* '03, i 1091.

SOEMNOFORM — Stated to be a mixture of Ethyl Bromide 1, Ethyl Chloride 12, and Methyl Chloride, 7, a rapid, safe and easily eliminated anæsthetic for use in dentistry — *L* '03, ii 635, *P.J.* '03, i 872.

Two fatal cases have been recorded in connection with Soemnoform, and it is stated (*L* '04, ii 1408) that, although the mixture was employed for the object of testing its effect, its use was not likely to be repeated.

Kelene is a proprietary article stated to consist of pure Ethyl Chloride, and put up in tubes fitted with patented automatic stopper. Useful for producing local anæsthesia. For general anæsthesia it is put up in graduated tubes.

Anæsthy and **Coryl** are stated to be mixtures of Ethyl and Methyl Chlorides.

Not Official **ÆTHYL IODIDUM**

ETHYL IODIDE HYDRIODIC FLUID

C₂H₅I, eq 154.72

A colourless, volatile, heavy and non inflammable liquid, with an agreeable ethereal odour and pungent taste.

It is produced by the action of resublimed Iodine on pure Ethylic Alcohol in the presence of amorphous Phosphorus. The coloured product is freed from uncombined Iodine by shaking with a solution of Sodium Bisulphite and purified by digestion over fused Calcium Chloride and redistillation.

It should be kept in well stoppered glass bottles of a dark amber tint and in a cool place.

It soon acquires a reddish-brown colour on exposure to light, but if no deeper than a pale wine colour it may be disregarded

The change of colour can be prevented by putting in the bottle a globule of Mercury

Solutions which have already become discoloured may be shaken with some finely-powdered Sodium Bisulphite, filtered and redistilled

Solubility—1 in 440 of Water, mixes in all proportions with Alcohol (90 p.c.).

Medicinal Properties—Antispasmodic It is used as an inhalation; 15 to 20 drops inhaled through the nose from a wide-mouthed bottle is more accurate and economical than dropping it on a handkerchief It is said not to weaken the digestive organs but rather to have a tonic effect It has been inhaled with success to relieve the dyspnoea in chronic bronchitis and asthma, also in secondary and tertiary syphilis as an adjunct to the administration of Potassium Iodide, the Iodine being very rapidly absorbed—*Squibb*, *B M J* '89, 11 1216; *P J* (8) xix 46

Prescribing Note—Can be obtained in glass capsules, 5 minims - 0.3 c.c. in each

Foreign Pharmacopœias—Mex (Eter Yodhidrico) Not in the others

Tests—Sp. gr. at 15.5° C (60° F) 1.943 Boiling point 71° to 72° C (159.8 to 161.6° F) It should leave no residue upon evaporation When shaken with an equal volume of Water and a little fuming Nitric Acid free Iodine is liberated, recognised by the reddish-brown colour produced The aqueous layer remaining after shaking together equal volumes of Ethyl Iodide and Water shall yield no turbidity with Silver Nitrate Solution (absence of Hydriodic Acid)

Not Official.

AGARICUS ALBUS.

AGARIC OF THE LARCH WHITE OR PURGING AGARIC

A species of mushroom, found growing on old Larches in Southern and Central Europe

Medicinal Properties—Has been used with success in night sweating of phthisis, checking cough and promoting sleep, also in hæmoptysis It has a strong cathartic action—*P* xxix 321, *M T* '81, 11 442, *T G* '88, 41, 371.

Dose—5 to 30 grains = 0.32 to 2 grammes of the powder given in jam

Foreign Pharmacopœias—Official in Fr, Ital (Agarico Bianco), Mex (Agarico Blanco), Port (Agarico Branco), Span and Swiss Not in the others.

Descriptive Notes—This fungus is not a true agaric, but belongs to the *Polyporei* in which the gills are replaced by slender tubes, giving a porous appearance to the under surface or the cap or pileus The commercial article consists of the fleshy part or stroma of the fungus deprived of the brown ringed white cuticle of the upper surface and, almost entirely, of the short tubular portion or hymenium of the lower surface It grows on various species of Larch, from Central Europe to Siberia, chiefly on *Larix Sibirica*, Led., and comes to England *via* Hamburg It varies in size from 3 to 8 inches (7.5 to 20 cm.) or more in diameter It is whitish, spongy and friable Although normally of the form of a rounded cone, it is often broken up into irregular pieces The taste is sweetish at first and afterwards bitter and acid, but it has no characteristic odour It can be rubbed to powder in a sieve, but becomes flattened when pounded in a mortar The active principle Agaricin is official in the *P.G.* Under the microscope the drug is seen to consist of slender hyphal threads, mixed with minute calcareous concretions

AGARICIN (Agaricic Acid)—A white crystalline powder. Melts at 138° C. (280.4° F)

Solubility—1 in 140 of Alcohol (90 p.c.); practically insoluble in Water and in weak Alcohol

Dose — $\frac{1}{4}$ to $1\frac{1}{2}$ grains = 0.016 to 0.1 gramme Generally given with Dover's Powder in a pill

Ph Ger maximum single dose, 0.1 gramme

It should not be given hypodermically —*L M R* '84, 118

In pill form $\frac{1}{12}$ grain very successful in night sweats of phthisis —*T G*, '94, 627

Foreign Pharmacopœias —Official in Dan, Ger, Ital, Jap, Mex and Norw Not in the others

ALCOHOL ABSOLUTUM.

ABSOLUTE ALCOHOL

FR, ALCOOL EMPHYRIQUE, GER, ABSOLUTER ALKOHOL, ITAL, ALCOOL ASSOLUTO SPAN, ALCOHOL ANHIDRO

A clear colourless, mobile, hygroscopic liquid, volatile and inflammable. It possesses a spirituous odour and a burning taste, and contains Ethyl Hydroxide, C_2H_5OH , eq 45.70, with not more than 1 p.c., by weight, of Water. It is obtained by the dehydration and rectification of Ethylic Alcohol of weaker strengths. It is possible to rectify Alcohol up to 98 p.c., beyond this dehydrating agents are necessary.

On account of its hygroscopic nature and its inflammability it should be preserved in well-stoppered bottles in a cool place.

Foreign Pharmacopœias —Official in Ital, sp gr 0.800, Span, sp gr 0.794, Dutch, sp gr 0.794 to 0.799, Austr, Belg, Dan, Ger and Jap, sp gr 0.796 to 0.800, Fr, sp gr 0.79433, Swed, sp gr 0.7955 to 0.8005, Swiss, sp gr not higher than 0.796, Mex (Alcohol Vinico), sp gr 0.790, U.S., sp gr not higher than 0.797 at 15.6° C (60° F), or 0.790 at 25° C (77° F). Not in the others.

Official Preparations —Used in the preparation of Chloroform, Liquor Ethyl Nitritus, and Liquor Sodii Ethylatis.

ALCOHOL (90 p.c.) —This is described under the heading *Spiritus Rectificatus*, as are also the weaker strengths of Alcohol, which are prepared from it.

Tests —The distinguishing tests for Absolute Alcohol are the sp. gr. which should be about 0.794, and the boiling point 78.5° C (173.3° F), the latter is not included in *BP* or *USP*, the *PG* gives 78.5° C (173.3° F).

The official figure for the sp. gr. is 0.794 to 0.796, the *USP* gives not higher than 0.797 at 15.6° C (60° F), or 0.790 at 25° C (77° F), the *PG* 0.796 to 0.800.

The more generally occurring impurities are excess of Water, extractive matter, substances of an oily or resinous nature, Amylic Alcohol, Fusel Oil, organic impurities, Aldehyde, and Tannic Acid. Absolute Alcohol is required to be neutral to blue Litmus paper by *US* and *PG*, but *BP* makes no mention of its reaction.

BP uses 1 to 2 p.c. of anhydrous Copper Sulphate as a test for excess of Water, requiring that when mixed with this quantity of the reagent and shaken occasionally within 2 or 3 hours the salt shall not become decidedly blue, but *USP* relies upon the sp. gr. alone.

All three Pharmacopœias agree that no fixed residue shall be left

upon evaporation, and that it shall mix with Water without any turbidity, indicating the absence of oily and resinous substances.

The *BP* and *USP* require that it shall leave no unpleasant or foreign odour when allowed to evaporate from clean filter paper, indicating the absence of Fusel Oil and allied impurities, the *PG* states that the Alcohol itself should possess no foreign odour.

The *BP* and *USP* differ in their method of testing for Amylic Alcohol, and in the quantities used for the test. *BP* indicates 100 c.c. of Absolute Alcohol and 2 c.c. of Volumetric Silver Nitrate Solution, decanting the supernatant liquid from the black precipitate formed during the first 24 hours' exposure to bright light, and requiring that the liquid shall undergo no further change when exposed to the light after the addition of more Volumetric Silver Nitrate Solution, *USP* employs 20 c.c. of Absolute Alcohol for the test and 1 c.c. of Silver Nitrate Test-solution, stipulating that no more than a faint opalescence nor more than a faint brownish tint shall be acquired when exposed for six hours to diffused daylight. The *PG* employs 10 c.c. of the Alcohol and 5 drops of Silver Nitrate Solution, requiring that it shall neither become turbid nor coloured on warming. The *PG* includes a test for Fusel Oil by evaporating a mixture of 10 c.c. of Absolute Alcohol and 0.2 c.c. Potassium Hydroxide Solution (15 p.c. w/w) to one-tenth its volume, and supersaturating the residue with Sulphuric Acid; when no odour of Fusel Oil should be developed.

The *PG* performs the test for readily carbonizable organic impurities on the Absolute Alcohol direct, without evaporation. The *USP* directs the spontaneous evaporation of the Alcohol (carefully protecting the liquid from dust during the evaporation) and the application of a few drops of colourless Sulphuric Acid to the residue. Both Pharmacopœias require that no red coloration should be produced.

USP and *PG* give practically the same test for limit of Aldehyde.

The Ammonia test for Tannic Acid and excess of Aldehyde is common to *BP* and *PG*, but is omitted from *USP*. The *BP* requires that no immediate coloration shall be produced on the addition of Ammonia Solution, the *PG* that no coloration shall be yielded on the addition of Ammonia Solution.

The *USP* includes a test for the absence of not more than 2 p.c. of Methyl Alcohol which does not appear in either *BP* or *PG*. It depends upon the oxidation of the Methyl Alcohol by means of Copper wire, and a test for Formaldehyde in the oxidised liquid by the Resorcin test. The Acetaldehyde produced by the oxidation of the Ethylic Alcohol is previously removed by boiling. The test is given in the small type under Spiritus Rectificatus, under the heading of Copper Wire and Resorcin.

The *PG* requires that the red coloration produced in a mixture of 10 c.c. of Absolute Alcohol and 1 c.c. of Potassium Permanganate Solution (0.1 p.c. w/w), shall not change to yellow within 20 minutes, the test is intended to detect the presence of Aldehyde or Tannic Acid, Fusel Oil and other organic impurities. This latter Pharmacopœia also requires that Absolute Alcohol shall not be

coloured by Hydrogen Sulphide Solution, indicating the absence of Copper and Lead

Except so far as strength is concerned, both *BP* and *USP* require Absolute Alcohol to respond to the respective tests given under 'Spiritus Rectificatus'

Not Official

ALCOHOL METHYLICUM

METHYLIC ALCOHOL

Syn—RECTIFIED PYROXYLIC SPIRIT

A colourless liquid, with a peculiar spirituous odour, and which has been submitted to various processes of rectification. It is produced by the destructive distillation of wood.

Its vapour forms explosive mixtures with air, and as it is very volatile it should be kept in well stoppered bottles and in a cool atmosphere.

Solubility—It mixes readily with Water, Ethylic Alcohol, Chloroform, and Ether. It dissolves Fats and volatile Oils.

Medicinal Properties—Narcotic, sedative, and anti emetic. It palliates the cough and lessens the febrile excitement of phthisis. It has been mixed with Chloroform for use as an anæsthetic (Regnault's Anæsthetic Mixture). *See CHLOROFORM*

In cases of poisoning, the use of the stomach pump and rectal injections are recommended (*BMJ* '05, i 262) to get rid of the poison. Stimulants and application of heat to the body and extremities. The treatment of the amaurosis is unsatisfactory. In the early stages, Pilocarpine and Potassium Iodide are indicated, and in the later stages Strychnine hypodermically or by the mouth.

The deodorised product has received considerable attention on account of its poisonous properties. Injected in small and continuous doses, its effect is far more deadly than grain Alcohol. The fumes are a distinct menace to eyesight and general health, and its external use is also objected to.—*L* '04, ii 1255

Tests.—Sp gr about 0.803. Boiling point about 55° to 66.5° C (131° to 151.7° F). It should be without action on Litmus paper, and should not be rendered turbid by admixture with Water. It should leave no empyreumatic odour on evaporation, and should be free from fixed residue. It should yield little or no reaction for Iodoform when tested with Potassium Hydroxide Solution and Iodine (absence of Acetone).

Dose—5 to 10 minims = 0.3 to 0.6 cc

Wood Spirit, Wood Naphtha, Pyroxylic Spirit are names applied to the crude article of commerce, which may contain from 75 to 90 p.c. of real Methylic Alcohol.

METHYLATED SPIRIT—*See SPIRITUS METHYLATUS*

METHYLIC ETHER—It is gaseous at ordinary temperatures, but is condensed by cold and pressure to a liquid boiling at -20° C (-4° F). A solution of this in Ethylic Ether is useful for producing local anæsthesia.

Not Official

ALETRIS

STAR GRASS COLIC ROOT

A perennial plant indigenous to U.S. The root was formerly included in the U.S. secondary list. It is stated to be useful as a uterine tonic, and has been employed with asserted benefit in colic, dropsy, and in chronic rheumatism.

Descriptive Notes—The rhizome occurs in pieces about 1 to 2 inches (25 to 50 mm) long, rarely branched, $\frac{1}{4}$ to $\frac{1}{2}$ inch (4 to 12.5 mm) in diameter,
1 2

impressed, covered with the brownish-grey scaly remains of leaves, and root-fibres, so that its outer surface is hidden. The transverse section is yellowish-white, spongy and porous, exhibiting here and there glistening points. The taste is mucilaginous and then bitter. It contains much starch, and unless kept in a bottle it is liable to be attacked by insects. It is derived from *Aletris* (Cannoe), L. and belongs to the nat. ord. Hamodoraceæ.

EXTRACTUM ALETRIDIS LIQUIDUM—A 1 in 1 fluid extract of the root prepared by percolation. The Alcohol is removed by distillation from the greater portion of the percolate, the residue is dissolved in the reserved portion, and sufficient of the menstruum (Alcohol 45 p.c.) added to produce the required volume. *B.P.C. Formulary* 1901.

This has been incorporated in the *B.P.C.*

ELIXIR ALETRIDIS.—Fluid Extract of Aletridis, 1; Fluid Extract of Liquorice, $\frac{1}{2}$; Tincture of Orange, $\frac{1}{2}$; Syrup, $1\frac{1}{2}$; Distilled Water, to yield 4.—*B.P.C. Formulary* 1901.

Dose.—30 to 60 minims = 1.8 to 3.6 c.c.

Elixir Aletridis Liquid Extract of Aletris, 25; Liquid Extract of Liquorice, 6; Simple Elixir, 45; Distilled Water, q.s. to produce 100.—*B.P.C.*

ALOES.

FR., ALOES; GER., ALOE, ITAL., ALOE, SPAN., ACIBAR

Both Aloe Barbadosensis and Aloe Socotrina are official in *B.P.* and *U.S.P.* *P.G.* has the African varieties only. See below.

Medicinal Properties.—Bitter tonic, purgative, acting chiefly on the large intestine, the slowest of purgatives, taking ten to fifteen hours to act. Stomachic bitter in very small doses. A good tonic cathartic in habitual constipation and in that associated with amenorrhœa and anæmia. Emmenagogue; should not be given during advanced pregnancy nor in inflammatory conditions of the pelvic organs. Small doses relieve, large doses aggravate hæmorrhoids. Used as an enema it is anthelmintic.

The aqueous extract is more active than is the resinous portion of Aloes, and the Barbados Aloes, containing a larger amount of aqueous extract than the Socotrine, is the more powerful; thus, 2 grains are equal to 3 grains of Socotrine.

Dose.—2 to 5 grains = 0.13 to 0.32 gramme.

Prescribing Notes.—Can be made into pills with a small quantity of diluted Alcohol; rarely prescribed alone.

1 grain with $\frac{1}{2}$ grain Extract of Nux Vom. is an excellent pill to obtain the stomachic effect, and to relieve habitual constipation. The *Pilula Aloes et Ferri*, and *Pilula Aloes et Myrrhæ* are given in amenorrhœa associated with chronic dyspepsia and constipation.

Official Preparations.—Of Barbados Aloes, *Extractum Aloes Barbadosensis*, *Pilula Aloes Barbadosensis*, *Pilula Aloes et Ferri*. Contained in *Pilula Cambogiæ Composita*, *Pilula Colocyntidis Composita*, and *Pilula Colocyntidis et Hyoscyami*. Used in the preparation of *Aloumum*. Of the **Extract**, *Decoctum Aloes Compositum*, *Extractum Colocyntidis Compositum*, *Tinctura Aloes*. Of **Socotrine Aloes**, *Pilula Aloes et Asafetidæ*, *Pilula Aloes et Myrrhæ*, *Pilula Aloes Socotrinæ*. Contained in *Pilula Rhei Composita*, *Tinctura Benzoini Composita*. Also used in the preparation of *Aloumum*.

Not Official.—Aloe Capensis, *Decoctum Aloes Compositum* 'Squire,' *Pilula Aloes Diluta*, *Pilula Aloes et Belladonnæ*, *Pilula Aloes et*

Nucis Vomicae, Pilula Aloes Composita, Pilula Aloes et Myrrhae, Pilula Aperientes Stahlii, Pilula Guttæ Aloeticæ, Pilula Laxativæ Composita, Pulvis Aloes et Canellæ, Tinctura Aloes Composita, Tinctura Aloes et Myrrhae, and Vinum Aloes

The distinguishing tests for Barbados and Socotrine Aloes will be found under their official headings. The African varieties are official in the *P G*, and the Descriptive Notes and Tests are given here.

South African Aloes is the evaporated juice of the leaves of the African varieties of the genus 'Aloe,' and forms a dark brown mass possessing a characteristic odour and bitter taste. It easily breaks into large pieces with a conchoidal fracture and sharp edged reddish splinters which are translucent and of a reddish or light brown tint, but do not possess any crystalline character under the microscope.

Descriptive Notes—There are three forms of Aloes imported from British South Africa. First, the vitreous blackish Aloes with a conchoidal fracture, and greenish brown translucent splinters, commonly known in this country as Cape Aloes, which give the reactions described in the *P G*. Second, an opaque or 'Hepatic' Cape Aloes, giving an orange brown powder, imported of recent years from Mossel Bay under the name of 'Uganda' Aloes, and manufactured from the leaves of *Aloe ferox*, L., and dried by sun heat alone, in the form of square bricks about four inches square and one inch thick. Third, an opaque Aloes very brittle and giving a pale greenish brown powder. It is rarely imported, and comes from Port Natal, but is remarkable for giving a permanent crimson colour with Nitric Acid and a bluish colour when Sulphuric Acid is added to it and the vapour of Nitric Acid blown over the mixture. It should perhaps be called Hepatic Natal Aloes, since a translucent Aloes is also prepared from *Aloe platy-lepis*, J. G. Baker, near Pietermaritzburg in Natal, and usually passes in commerce for Cape Aloes and gives the same chemical reaction as that prepared from *Aloe ferox*, L. The botanical source of Hepatic Natal Aloes is unknown.

Tests—The distinguishing test for African Aloes is that if a particle be treated with Nitric Acid no red but only a pale greenish zone shall be formed within three minutes (differing thus from Barbados Aloes).

About 40 p.c. should be soluble in cold Water, and it should be completely soluble 1 in 5 (by weight) of warm Alcohol (90 p.c.), the solution remaining clear even after cooling.

5 parts of Aloes treated with 60 parts of boiling Water should yield an almost clear solution, from which about 3 parts again separate on cooling, *P G*.

Aloes dissolved in hot Water produces with Concentrated Solution of Borax a greenish fluorescence, *P G*.

It should impart to boiling Chloroform and Ether only a very faint yellow colour, and Ether so coloured should leave only a very slight residue, *P G*.

Thoroughly dried Aloes is used for the preparation of Powdered Aloes. The powder should not agglutinate at 100° C (212° F) nor alter in colour, *P G*.

ALOE BARBADENSIS. BARBADOS ALOES

FR., ALOES DES BARBADES, GER., BARBADOS ALOE, ITAL., ALOE VFRA., SPAN., ACIBAR BARBADO.

They are obtained principally from the leaves of *Aloe vera*, L., and *Aloe chinensis*, Steud., in the West Indian Islands, and sold as Barbados or Curaçao Aloes.

Solubility.—Water dissolves about 75 p.c. It is almost entirely soluble in Alcohol (60 p.c.).

Dose.—2 to 5 grains = 0.13 to 0.32 gramme.

Foreign Pharmacopœias.—Official in Belg., Dutch, Fr., Ital., Jap., Port., Spain and U.S., Mex (Acibar). Not in the others.

Descriptive Notes.—Barbados or Curaçao Aloes as imported varies considerably in colour, consistence, and degree of opacity, and

may be in the form of a stiff yellowish-brown paste, a mass of pitchy character, or hard and solid, but it always has a comparatively smooth surface, when dried it varies in colour from chocolate brown to black, or rarely exhibiting more or less translucent pieces, which become opaque on keeping. It has a characteristic somewhat nauseous odour, more evident when freshly broken, or powdered, and a very bitter taste. The best qualities conform to the requirements as to solubility given under Tests. Such Aloes as present a chocolate brown colour and freedom from air vesicles are of the finest quality and should be chosen for medicinal use, those characters indicating evaporation of the fresh, not fermented, juice by steam heat, black vesicular samples being of inferior quality, prepared from fermented juice and evaporated over a naked fire.

Aloes are now rarely imported from Barbados, and then only in small quantity, coming almost exclusively from the Dutch West Indian Islands of Curaçao, Bonaire, and Aruba.

Tests.—The distinguishing test for Barbados Aloes is the crimson colour imparted to Nitric Acid by the powder.

The more generally occurring impurity is an undue amount of insoluble matter. This is guarded against by the requirement that not less than 70 p.c. shall be soluble in Water, and that it should be almost entirely soluble in a mixture containing Alcohol (90 p.c.) diluted with half its volume of Water.

Barbados Aloes yields about 2 p.c. of ash, and 3 p.c. should not be exceeded.

1 **Borntrager's test**—Shake out with Benzene, and treat separated Benzene with Ammonia, pink colour on standing.

2 **Klunge's test**—Warm with Copper Sulphate and a little Sodium Chloride, yellow colour, changing to red or violet. Fair test for West Indian Aloes, but not much good for the other varieties.

3 **Flückiger's test**—Sulphuric Acid and Nitric Acid Vapour, deep blue colour. Specific test for Natal Aloes.

3 **Bainbridge's test**—Nitric Acid, red colour, changing to green. Distinguishes it from Cape Aloes.

5 **Cripp's and Dymond's test**.—Triturate 1 grain of sample with 16 drops Sulphuric Acid and 4 drops Nitric Acid and dilute with an oz. of Water. A deep orange to crimson colour is developed, intensified by the addition of Ammonia. This appears to be the best general test for Aloes.—*J. J.* (8) xv 633. The reaction is also given by all bodies containing or yielding Chrysanthemum Acid, but these yield a pink colour with Ammonia alone while Aloes gives a yellow.

ALOE SOCOTRINA. SOCOTRINE ALOES

FR, ALOES SOCOTRIN, GER, SOCOTRINISCHE ALOE, ITAL, ALOE DI SOCOTORA; SPAN, ACIBAR SOCOTRINO

Socotrine or Zanzibar Aloes, obtained from *Aloe Perryi*, J. G. Baker, and possibly other species. Imported principally through Bombay.

Solubility.—Water dissolves about 50 p.c.; the residue is pretty well inert, almost entirely soluble in Alcohol (60 p.c.)

Dose.—2 to 5 grains = 0.13 to 0.32 gramme

Foreign Pharmacopœias—Official in Belg., Ital., Mex., Port., Spain and U.S., U.S. has also Aloe Purificata, which is Socotrine Aloe dissolved in Alcohol, strained and evaporated to dryness. Cape Aloe is official in Austria, Belg., Dan., Dutch, Fl., Ger., Hung., Ital., Norw., Port., Russ., Spain, Sweden and Swiss.

Descriptive Notes—Socotrine Aloe is usually imported in barrels, in a semi-liquid or pasty condition, and sometimes in a fermented or more or less fetid state. When dried it presents an extractiform or irregular surface, is opaque, and if of good quality is of a dark brown colour and gives an orange-brown powder. It possesses a not unpleasant odour. Inferior qualities are black and vesicular and have a rancid or butyric odour, and are unfit for medicinal use. Zanzibar Aloe is imported in skins, containing several pounds, packed in rectangular cases. It is usually of better quality than the Socotrine Aloe, and is commonly sold as 'Hepatic' Aloe, the opaque fractured surface being of a liver colour. Samples of which the splinters are garnet red and translucent are now rarely met with in either the Socotrine or Zanzibar varieties. These Aloes should conform to the requirements as to solubility given under Tests, and also the reactions with Nitric Acid. An East African Aloe met with in commerce is sold as Socotrine which does give a crimson colour with Nitric Acid, and therefore is excluded by the *B P* test, whilst its geographical source prevents its inclusion under Barbados Aloe; this kind exhales a faint odour like that of Curaçao Aloe, when freshly broken or powdered. There is also an Aloe, which comes *via* Bombay, from Jafferabad and other parts on the Arabian coasts, which does not give a crimson colour with Nitric Acid, but this kind yields a powder of a dull brown, not of an orange-brown colour like the Zanzibar and Socotrine Aloes, and possesses hardly any odour. The official description does not exclude Jafferabad Aloe, except in so far as that name is not used.

Tests—The distinguishing test for Socotrine Aloe is the reddish or yellowish-brown colour produced when the powder is treated with a drop or two of Nitric Acid.

The more generally occurring impurities are leaves of *Calotropis* and stones, and an undue proportion of insoluble matter, but the opaque Natal Aloe might be mistaken for it, although its powder has a greenish-brown hue, not the orange-brown tint of Socotrine and Hepatic Aloe. The distinction of Barbados Aloe from Natal Aloe is ensured by the official requirement that no blue coloration shall be produced when the vapour of Nitric Acid is blown over the powder, previously moistened with Sulphuric Acid, the absence of the other impurities by the fact that *B P* requires about 50 p.c. to be soluble in Water, and that it should be almost entirely soluble in a mixture of Alcohol (90 p.c.) with half its volume of Water. The *U S P* fixes a standard for moisture, which should not exceed 10 p.c.

Socotrine Aloe yields about 2 p.c. of ash, and 3 p.c. should not be exceeded.

ALONUM. ALOIN $C_{16}H_{16}O_7, 3H_2O$, eq 371.36

A yellow crystalline powder possessing a faint odour of Aloes and a very bitter taste. It is a neutral, non-glucosidal, bitter principle, obtained chiefly from Barbados Aloes.

It may be assumed that commercial 'Alon' is α -Barbaloin. Its formula is $C_{16}H_{16}O_7$, with about three molecules of Water of crystallisation.

Solubility—1 in 120 of Water, 1 in 18 of Alcohol (90 p.c.), freely soluble in hot Water, nearly insoluble in Ether.

B.P. states sparingly soluble in cold Water, *U.S.P.* gave the figure as 1 in about 65 at 25° C (77° F), but this was subsequently altered in the list of corrections and additions to 1 in 120, 3 samples obtained (1903) from different manufacturers yielded a solution at 1 in 120.

Dose.— $\frac{1}{2}$ to 2 grains = 0.03 to 0.13 gramme.

Prescribing Note—Generally given in pills or in cachets with other ingredients. 'Diluted Glucose' is a good excipient for Alon in pills.

Not Official—*Pilula Aloini Composita*, *Pilula Aloini et Podophylli*.

Foreign Pharmacopœias—Official in U.S. Not in the others.

Aloins may be classified as follows—

BARBALOINS—Yielding on oxidation Chrysammic, Aloetic, and Picric Acid.

α -barbaloin, which gives a red colour with cold Nitric Acid (1.42), obtained from Barbados and Curaçao Aloes.

β -barbaloin, which requires either fuming Nitric Acid, or a hot Acid of ordinary strength to give the red coloration. This variety is yielded by Jafferabad, and by some varieties of Socotrine and Zanzibar Aloes, see p. 119.

NATALOIN—Yields on oxidation Picric but not Chrysammic Acid. This is a distinct species, from Natal Aloes only, having a formula $C_{24}H_{20}O_{10} \cdot H_2O$. Softens at 180° C., and melts at 210° C.

Tests.—The distinguishing tests for Barbaloin are its melting point, which, when anhydrous, should be 147° C (296.6° F), and the red coloration produced on the addition of Nitric Acid. Both *U.S.P.* and *B.P.* state that Alon is rapidly affected in alkaline solutions, but only slowly in neutral or acidified solutions.

B.P. gives no tests for impurities, but those more generally occurring are mineral residue, Aloins derived from Natal or Cape Aloes, and Emodin. Mineral matter is detected by the ash left on ignition, whilst Emodin is detected by treating the 10 p.c. Benzene Solution with an equal volume of 5 p.c. Ammonia Water. It should leave no weighable residue when ignited with free access of air. In testing for Emodin a weighed quantity of Alon is shaken with 10 c.c. of Benzene for 1 minute and filtered, the filtrate should not impart more than a faint pink colour to an equal volume of Ammonia Solution (5 p.c.) when shaken with it. Its distinction from Nataloin, Socaloin and Capaloin is carried out by the colour tests mentioned below.

Colour reactions of the *U.S.P.* (a) The solutions in Ammonia and the Alkalis are yellow, turning red with a greenish-red fluorescence. (b) Nitric Acid gives with Curaçao Alon a cherry-red solution (distinction from Nat. Soc.

and Capaloin) (c) Sulphuric Acid with a minute quantity of Aloin forms a yellowish-red solution, which with a small crystal of Potassium Dichromate added becomes olive green, then dark green, and finally blue on standing, with a larger amount of Potassium Dichromate, the yellowish red solution first turns purple, then brown, and finally green (d) Bromine Water colours an aqueous solution pink (e) Gold Chloride T S to carmine-red, turning to violet (f) Ferric Chloride T S with an alcoholic solution gives a brownish green colour (g) Copper Sulphate T S gives with a dilute aqueous solution of Curaçao Aloin a bright yellow colour, this mixture with a few drops concentrated solution of Sodium Chloride gives a red colour, and on further adding a little Alcohol the colour becomes violet (distinction from Nataloin and Capaloin)

Preparations

DECOCTUM ALOES COMPOSITUM COMPOUND DECOCTION OF ALOES *N O Syn* —BAUME DE VIE

An aqueous solution prepared by boiling together for five minutes Extract of Barbados Aloes 1, Myrrh $\frac{1}{2}$, Potassium Carbonate $\frac{1}{2}$, Extract of Liquorice 4, and Distilled Water 40 Immediately after the boiling is finished add Saffron $\frac{1}{2}$, and, when the liquid has cooled down, Compound Tincture of Cardamoms 30, and more Distilled Water to make 100 of product (1 of Extract in 100)

Dose — $\frac{1}{2}$ to 2 fl oz = 14.2 to 56.8 c c

Tests —Decoction of Aloes, *B P*, has a specific gravity of about 1.005, contains about 5.5 p c w/v of total solids and about 20 p c w/v of Absolute Alcohol

DECOCTUM ALOES COMPOSITUM 'SQUIRE' Made with Socotrine Aloes and the Fluid Extract of Liquorice

Dose — $\frac{1}{2}$ to 2 fl oz = 14.2 to 56.8 c c

The fluid extract is much better than the solid extract for covering the taste of Aloes, there is a marked difference in the taste of the two preparations, even when they practically contain the same amount of Liquorice This suggestion has been adopted in *B P* in the case of Tincture of Aloes, but not in that of the Compound Decoction where it is of more importance

Tests —Squire's Decoction of Aloes has a specific gravity of about 1.009, it contains about 10 p c w/v of total solids and about 22 p c w/v of Absolute Alcohol

EXTRACTUM ALOES BARBADENSIS.—EXTRACT OF BARBADOS ALOES

An aqueous Extract, of which about $\frac{3}{4}$ grain is equal to 1 grain of the Aloes

Dose.—1 to 4 grains = 0.06 to 0.26 gramme

Foreign Pharmacopœias —Extract of Aloes is Official in Austr, Dan, Dutch, Ger, Hung, Ital, Jap, Norw, Russ, Swed, Swiss and U.S Not in the others

PILULA ALOES BARBADENSIS PILL OF BARBADOS ALOES.

4 of the pill is about equal to 2 of Barbados Aloes, 1 of Hard Soap, $\frac{1}{2}$ of Oil of Caraway, and 1 of Confection of Roses

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

PILULA ALOES SOCOTRINÆ. PILL OF SOCOTRINE ALOES.

4 of the pill is about equal to 2 of Socotrine Aloes, 1 of Hard Soap, $\frac{1}{8}$ of Oil of Nutmeg, and 1 of Confection of Roses

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

Foreign Pharmacopœias—Official in Fr (Pilules d'Aloes et de Savon), Jap and US (Pilulæ Aloes), Aloes and Soap, equal parts, Mex (Pildoras de Acibar), Aloes 10, Soap 2, Swiss (Pilula Aloetica), Aloes 10, Soap 1, Glycerin 8 drops, Spirit $q\ s$. Not in the others

PILULA ALOES ET ASAFETIDÆ. PILL OF ALOES AND ASAFETIDA

4 of the pill is about equal to 1 of Socotrine Aloes, 1 of Asafetida, 1 of Hard Soap, and 1 of Confection of Roses

Dose.—4 to 8 grains = 0.26 to 0.52 gramme.

Foreign Pharmacopœias—Official in US, 1 in 3, Jap, equal parts of Aloes, Asafetida, Soap and Honey. Not in the others

PILULA ALOES ET FERRI. PILL OF ALOES AND IRON.

$4\frac{1}{2}$ of the pill is about equal to 1 of Barbados Aloes, $\frac{1}{2}$ of Exsiccated Ferrous Sulphate, $1\frac{1}{2}$ of Compound Powder of Cinnamon, and $1\frac{1}{2}$ (by weight) of Syrup of Glucose

Dose—4 to 8 grains = 0.26 to 0.52 gramme.

Foreign Pharmacopœias—Official in US, Purified Socotrine Aloes 1, Exsiccated Ferrous Sulphate 1, Aromatic Powder 1, Confection of Roses $q\ s$, Ger, Cape Aloes 1, Exsiccated Ferrous Sulphate 1, Sp Saponis $q\ s$, Jap, equal parts of Aloes and Spirit $q\ s$, Swiss, Aloes 5, Ferrous Sulphate 5, Soap 1, Glycerin 5 drops, Alcohol $q\ s$. Not in the others

PILULA ALOES ET MYRRHÆ. PILL OF ALOES AND MYRRH.

$4\frac{1}{2}$ of the pill is about equal to 2 of Socotrine Aloes, 1 of Myrrh, and $1\frac{1}{2}$ (by weight) of Syrup of Glucose

The composition of this pill, known also as Pil Rufi, remained much the same for about 300 years, but in 1898 the *BP* omitted the Saffron, the proportions of Aloes and Myrrh remaining the same.

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

Foreign Pharmacopœias—Official in Port and US, with Aromatic Powder in place of Saffron, Purified Aloes 13, Myrrh 6, Aromatic Powder 4, in grammes, Syrup $q\ s$ to make 100 pills. Not in the others

TINCTURA ALOES. TINCTURE OF ALOES

Extract of Aloes, 1, Liquid Extract of Liquorice, 6, Alcohol (45 p.c.), $q\ s$ to make 40 (1 Extract in 40)

Dose.— $1\frac{1}{2}$ to 2 fl drms = 5.4 to 7.1 c.c., when repeated, $\frac{1}{2}$ to 1 fl. drms = 1.8 to 3.6 c.c.

Foreign Pharmacopœias—Official in Belg, 1 in 5; Dutch, Fr., Ger, Ital, Jap, Russ, Span, Swiss, 1 and 5, Hung. and Port, 15 in 100, US, 1 in 10. All are by weight, except US

Tests.—Tincture of Aloes has a sp. gr. of 0.970 to 0.980; it contains from 7 to 8 p.c. w/v of total solids and about 40 p.c. w/v of Absolute Alcohol

Not Official

ALOE CAPENSIS (Cape Aloe) —A translucent variety See p 117

Official in all the Foreign Pharmacopœias

DECOCTUM ALOES COMPOSITUM 'SQUIRE'—See p 121

ENEMA ALOES —Aloes 40 grains, Carbonate of Potassium 15 grains, Mucilage of Starch 10 fl oz —*B P* 1885

Aloes 0.75, Potassium Carbonate 0.25, Glycerin 10, Mucilage of Starch *q s* to produce 100 — *B P C*

PILULA ALOES DILUTA —Marshall Hall's Pill Barbados Aloe 4, dissolve in Water and strain, then add Extract of Liquorice 4, Treacle 4, thinly sliced Hard Soap 4, mix and evaporate to a pilular consistence

Dose —3 or 4 grains = 0.2 or 0.26 gramme

This has been incorporated in the *B P C* as a 4-grain pill

PILULA ALOES ET BELLADONNÆ —Extract of Aloe, 1 grain, Extract of Belladonna, $\frac{1}{2}$ grain

PILULA ALOES ET NUCIS VOMICÆ —Extract of Aloe, 1 grain; Extract of Nux Vomica, $\frac{1}{2}$ grain

Barbados Aloe, 2 grains, Extract of Nux Vomica, $\frac{1}{2}$ grain; Alcoholic Extract of Belladonna, $\frac{1}{2}$ grain, in each pill —*St Thomas's*

This has been incorporated in the *B P C*, using Alcohol (60 p c) as an excipient

PILULA ALOINI COMPOSITA —Aloin, Extract Nucis Vomice, Ferri Sulphatis, Pulv Myrrhæ, Saponis, ana $\frac{1}{2}$ grain —*L '87*, 1.2 (Sir Andrew Clark's Laver Pill)

This has been incorporated in the *B P C*

PILULÆ APERIENTES STAHLII (*Swed*) —Extract Aloe, 6, Extract Rhei Co., 3, Reduced Iron, 2, Rad Althææ, 2, Alcohol (64 p c) and Simple Syrup, *q s* to make 100 pills

PILULÆ ALOES COMPOSITÆ *Syn* BAIRD'S PILLS —Barbados Aloe, in powder, 30, Ipecacuanha Root, in powder, 6, Scammony, 30, Green Extract of Hyoscyamus, 30, Syrup of Glucose, *q s*, in 100 parts. Mix to form a mass and divide into pills weighing 4 grains each —*P J F* and *B P C*

Dr Mair's Pills —Ipecacuanha Powder, 25 grains, Scammony, in powder, 2 drms, Extract of Aloe, 2 drms, Extract of Hyoscyamus, 2 drms, make a mass and divide into 5-grain pills —*Pharm Form*

Aloes Pilulæ Composita —*Belg*, Aloe 10, Scammony 3, Jalap 3, Ginger 4, Soap 10, *Ital*, Aloe 3, Jalap 3, Soap 3

Pilulæ Aloes et Jalap —Equal parts of Aloe, Jalap, Soap and Liquorice —*Jap*

PILULÆ ALOES ET MASTICHES —Purified Aloe, in fine powder, 13 grammes, Mastic, in fine powder, 4 grammes, Red Rose, in powder, 3 grammes, Alcohol (49 p c), *q s* to make 100 pills —*U S P*

Each pill will weigh about 3 grains. They are in imitation of Lady Webster's Dinner Pills, and one of them may be given as a laxative at bedtime or before a meal

Barbados Aloe, in powder, 65, Mastic, in powder, 20, Confection of Roses, 15, in 100 parts. Divide into pills weighing 4 grains each —*B P C*

In Gray's Supplement these pills are given as 3 grains each

PILULA ALOINI ET PODOPHYLLI COMPOSITA —Aloin 2, Capsicin 1, Jalapin 2, Podophyllum Resin 4, Green Extract of Hyoscyamus 1, Extract of Nux Vomica 1, dose of the mass, $\frac{1}{2}$ to 2 grains —*P J F*

Aloin 2, Oleoresin of Capsicum 1, Jalap Resin 2, Podophyllum Resin 3, Extract of Nux Vomica 1, Green Extract of Hyoscyamus 1; in 10 parts. Divide into pills weighing $\frac{1}{2}$ grain each —*B P C*

Little Antibilious Pills. —Podophyllum, 8 grains, Aloin, 6 grains, Jalapin, 6 grains, Capsicin, 8 grains, Ipecacuanha Powder, 3 grains, Extract of

Hvoscjarius 3 grains Extract of Nux Vomica, $2\frac{1}{2}$ grains; Glycerin Tragacanth, $q\ s$ to make a mass. Divide into 60 pills — *Pharm Form*

PILULÆ LAXATIVÆ COMPOSITÆ — Aloin, 1 3, Strychnine, 0 05, Extract of Belladonna Leaves, 0 8, Ipecac, in powder, 0 4, Glycyrrhiza, 4 6, in grammes, Syrup, $q\ s$ to make 100 pills — *U S P*

A modification of this has been incorporated in the *B P C* as follows —

Pilulæ Aloini et Strychninæ Compositæ — Aloin, 50, Strychnine, 5, Green Extract of Belladonna 25, Ipecacuanha, 12 50, Milk Sugar, $q\ s$, Syrup of Glucose, $q\ s$, in 100 parts, divide into $\frac{1}{2}$ -grain pills — *B P C*

PIL GUTTÆ ALOETICÆ (*Swed*) — Aloes, 7; Camboe, 3, Gum Arabic, 3, Galbanum, 4, Carvone, 1 5, Syrup, $q\ s$ to make 100

PULVIS ALOES ET CANELLÆ (*Hiera Picta*) — Powdered Socotrine Aloes, 4, Powdered Canella Bark, 1

This has been incorporated in the *B P C*.

TINCTURA ALOES COMPOSITA *Syn* ELIXIR AD LONGAM VITAM.—
Ger — Aloes, in coarse powder, 3, Gentian, cut middling fine, 0 5; Rhubarb, cut middling fine, 0 5, Zedoary, cut middling fine, 0 5; Saffron, 0 5, Alcohol (70 p c), 100

This has been incorporated in the *B P C*, but with the Alcohol by volume instead of by weight

Austr — Aloes, 30, Gentian, 5, Rhubarb, 5, Zedoary, 5, Saffron, 5; Alcohol, (68 p c), 1000

Fr — Aloes, 25, Agaric, 2 5, Gentian, 2 5, Rhubarb, 2 5, Zedoary, 2 5, Saffron, 2 5, Alcohol (60 p c), 1000.

Ger and *Jap* — Aloes, 30, Gentian, 5, Rhubarb, 5; Zedoary, 5; Saffron, 5, Alcohol (68 p c), 1000

Mex — Aloes, 8, Gentian, 1, Rhubarb, 1, Saffron, 1, Agaric, 1, Treacle, 1, Alcohol (60 p c), 400

Russ — Aloes, 45, Gentian, 5, Rhubarb, 5, Saffron, 5, Alcohol (70 p c), 1000.

Swiss — Aloes, 6, Saffron, 1, Agaric, 1, Myrrh, 1, Gentian, 1, Rhubarb, 1; Zedoary, 1, Alcohol (68 p c), 200

All are by weight

TINCTURA ALOES ET MYRRHÆ — Purified Aloes, 1, Myrrh, 1, Powdered Liquorice Root, 1, Alcohol (94 p c), 75, and Water, 25, mixed $q\ s$ to make 10 — *U S P*

Socotrine Aloes, in powder, 10, Saffron, 5, Tincture of Myrrh, 100 — *Edin Ph* 1841 *Syn* Elixir Proprietatis — *PL* 1721, *B P C*

VINUM ALOES — Socotrine Aloes, $1\frac{1}{2}$ oz, Cardamom Seeds, bruised, 80 grains, Ginger, in coarse powder, 80 grains, Sherry, 2 pints — *B P* 1835

This has been incorporated in the *B P C* as follows —

Socotrine Aloes, crushed, 3 75, Cardamom Seeds, bruised, 0 50, Ginger, in coarse powder, 0 50, Sherry, sufficient to produce 100

Not Official

ALSTONIA.

The dried bark of *Alstonia Scholaris*, R Br, and of *Alstonia Constricta*, F Mull Infusum Alstoniæ (1 in 20), dose $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c.c., and Tincture Alstoniæ (1 in 8), dose 30 to 60 minims = 1 8 to 3 6 c.c., are official in *Ind* and *Col Add.* for India and the Australian and Eastern Colonies.

Not Official.

ALTHÆÆ RADIX.

MARSHMALLOW ROOT.

Fr, GUIMAUME, *Ger*, EIBISCHWURZEL, *Ital*, ALTEA; *Span*, ALTEA.

Descriptive Notes — The root of the *Althæa officinalis*, Linn. Marsh-mallow root occurs in commerce in two forms, viz — (1) The natural root dried,

and (2) the root with the outer bark removed. The former is usually sold by herbalists, the latter by pharmacists. The natural root bears some resemblance to Liquorice root, in its external characters, the surface having short transverse scars, the bark being tough and finely fibrous. It is, however, white internally (Liquorice is yellowish-white) and finely fibrous and has a mucilaginous instead of a sweet taste. The decorticated root is often larger than the natural root, and deeply grooved longitudinally. The larger decorticated roots are probably often derived from the allied species *A. Narbonensis*, L. Both are very mucilaginous. It is necessary to preserve the root in a dry place, or the syrup prepared from it will turn yellowish and have a disagreeable odour. Lozenges made of the powdered root with gum and flavoured with orange flower water are sold as *Pastilles de Gumaueve*. The powder is also used in pill masses.

Medicinal Properties—It is much employed on the Continent as a demulcent in irritation and inflammation of the mucous membranes of the mouth and pharynx.

Official in all the Foreign Pharmacopœias

The two substances **Asparagin** and **Betain** have been extracted from *Althæa* root.

Asparagin dissolves Mercuric Oxide, but the Oxide must be freshly precipitated. A solution is best prepared by precipitating Mercuric Chloride Solution, washing the precipitated Oxide and dissolving it in solution of **Asparagin**. The solution has been employed as a hypodermic injection in the treatment of syphilis.

DECOCTUM ALTHÆÆ—*Althæa* Root, 1, Water, 30, boil to 20. This has been incorporated in the *B P C*.

SYRUPUS ALTHÆÆ—Macerate 3 of *Althæa* Root in 40 of Water for twelve hours, strain, press, and filter until 32 have passed through, to this add 64 of Sugar, dissolve warm, and heat the Syrup to boiling, when cold, skim and strain through flannel.

This has been incorporated in the *B P C*.

Foreign Pharmacopœias—Official in all except Belg, Fr and U S.

TROCHISCI ALTHÆÆ—About 1 gram in each lozenge. Demulcent. Valuable after excision of tonsils or uvula.

Foreign Pharmacopœias.—Official in Ital, Mex. (*Pastillas de Altea*), Span. Not in the others.

ALUMEN.

ALUM

$\text{Al}_2(\text{SO}_4)_3, \text{K}_2\text{SO}_4, 24\text{H}_2\text{O}$, eq 941.94

$\text{Al}_2(\text{SO}_4)_3, (\text{NH}_4)_2\text{SO}_4, 24\text{H}_2\text{O}$, eq 900.16

FR, ALUN DE POTASSIUM, GER, KALIALAUN, ITAL, SOLFATO DI ALLUMINIO E DI POTASSIO, SPAN, SULFATO ALUMINICO-POTASSICO

Both salts are official, Potash Alum (Aluminium and Potassium Sulphate), and Ammonia Alum (Ammonium and Aluminium Sulphate). They are practically alike in appearance, occurring in large colourless octahedral crystalline masses, possessing a sweetish and very astringent taste.

Solubility. -1 in 11 of Water, 3 in 1 of boiling Water; Potash Alum, 1 in 3 of Glycerin, Ammonia Alum, 1 in 1½ of Glycerin. Insoluble in Alcohol (90 p.c.) *

Alum when heated melts in its own Water of crystallisation.

Medicinal Properties—Astringent, used as a gargle, mouth-wash, or spray for tonsillitis, aphthous conditions of the mouth, and pharyngitis, 10 grains in 1 oz of Water, as an injection in leucorrhœa and gonorrhœa, 60 grains in a pint of Water, as a nasal douche in chronic ozœna, 4 grains in 1 oz of Water, as a snuff in epistaxis, 6 grains mixed with 1 grain of Starch, as a lotion in purulent ophthalmia, 2 to 6 grains in 1 oz of Water. 10 to 15 grains three times a day have been given for internal hæmorrhage, such as that of typhoid or gastric ulcer, also for menorrhagia, and in cases of lead poisoning, arrests excessive secretion in dysentery, diarrhœa and night sweats; vomiting caused by the cough of phthisis is sometimes checked by 6 to 10-grain doses of Alum. A saturated solution of Water forms an excellent styptic for hæmorrhage of rectum, bleeding hæmorrhoids, epistaxis, etc., the glycerin of alum is used in inflamed tonsils. 60 grains have been recommended as an emetic in croup. Dried Alum is escharotic, used for warty growths and to stimulate indolent ulcers, and to destroy exuberant granulations and to remove nævi.

Dose—5 to 10 grains = 0.32 to 0.65 grammes

Prescribing Note—*Mostly used in aqueous or Glycerin solution*

Incompatibles.—Alkalis and their Carbonates, and Tannic Acid.

Official Preparations—Glycerinum Aluminis, Alumen Exsiccatum

Not Official—Alum Rose Gargle, Alumina, Aluminis, Aluminium Acetate Solution, Aluminium Aceto-Tartrate, Aluminium Chloride, Aluminium Nitrate, Aluminium Nitrate, Aluminium Oleate, and Aluminium Sulphate, Pessus Alu., and Zinc, Pulvis pro Pedibus

Foreign Pharmacopœias—Official in Austr., Bel., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex. (Sulfato de Aluminio y Potasio), Norw., Port., Russ., Span. (Sulfato Aluminico-Potasio), Swed., Swiss and U.S. All of them use Potash Alum only

Tests.—The distinguishing tests for Alum are that it shall yield, when dissolved in Water, a white gelatinous precipitate with Ammonia Solution or with Potassium or Sodium Hydroxide Solution, the white precipitate produced by Ammonia Solution is practically insoluble in excess of the reagents, whilst that produced by solution of Potassium or Sodium Hydroxide dissolves and is again reprecipitated by sufficient Ammonium Chloride Solution, indicating the presence of Aluminium. When boiled with Potassium or Sodium Hydroxide Solution it evolves a strong ammoniacal odour, and the issuing gas has a strongly alkaline reaction towards moistened red Litmus paper, indicating the presence of Ammonia, or a saturated aqueous solution yields with Tartaric Acid Solution or Sodium Bitartrate Solution a white crystalline precipitate within half an hour, indicating the presence of Potassium. It yields a white precipitate, insoluble in Hydrochloric Acid, on the addition of Barium Chloride Solution, indicating the presence of Sulphates.

The more generally occurring impurities are Calcium, Copper, Iron, and Lead.

The aqueous solution should, when faintly acidified with Hydrochloric Acid, be unaffected by Hydrogen Sulphide Solution, indicating

the absence of Copper and Lead, and when this solution is rendered alkaline by the addition of a sufficient excess of Ammonia Solution it should not materially darken in colour, indicating the absence of more than traces of Iron. The *P G* and *U S P* include a separate test for Iron which is given under the heading of Potassium Ferrocyanide Solution in the small type below, it should not afford a distinct turbidity on the addition of Ammonium Oxalate Solution, indicating the absence of more than traces of Calcium.

In *U S P* and *P G* only the Potash Alum is official. The *U S P* requires that it shall contain not less than 99.5 p.c. pure Aluminium Potassium Sulphate, and it shall lose 45.55 p.c. of its weight when all of its Water of crystallisation is driven off.

Potassium Ferrocyanide Solution—20 c.c. of a solution (1-20) should not be coloured blue immediately by 0.5 c.c. Potassium Ferrocyanide T.S., *P G* and *U S P*.

Preparations

GLYCERINUM ALUMINIS. GLYCERIN OF ALUM

Powdered Alum, 1 oz., Distilled Water, 3 fl. dm., Glycerin, *q s*
to make 6 oz. (1 in 6)

Pure Alum should and does dissolve clear in Glycerin, but commercial Pulv. Aluminis, as a general rule, will not dissolve without residue except after prolonged boiling.

A powerful local astringent. When diluted with Water it forms a useful gargle.

Sometimes prescribed with an equal quantity of Glycerin of Tannic Acid.

ALUMEN EXSICCATUM. EXSICCATED ALUM *Syn*—ALUMEN USTUM

Potash Alum, deprived of its Water by heat. It yields about 55 p.c. of product.

Foreign Pharmacopœias—Official in Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Norw., Port., Russ., Span., Swiss and U.S.

Not Official

ALUM ROSE GARGLE—Broken Rose petals, 3 drms., Diluted Sulphuric Acid, 3 fl. drms., cold Distilled Water, 10 fl. oz., digest for two hours, and strain 8 fl. oz., then add Alum, 2 drms., Sugar, 4 dm., Alcohol (90 p.c.), 4 fl. drms., dissolve. This kept well for seven years. To be mixed with an equal bulk of Water before use.

Gargarisma Aluminis—Alum, 2, Acid Infusion of Roses, *q s* to produce 100.—*B P C*

GOSSYPIUM ALUMINIS—Contains about 30 p.c. of Alum.

PESSUS ALUMINIS.—Alum, 15 grains, Oil of Theobroma, 2 drms.—*Westminster*

This has been incorporated in the *B P C*.

PESSUS ALUMINIS ET ZINCI—Exsiccated Alum, 5 grains, Zinc Sulphate, 5 grains, Opium, in powder, 1 grain, Basis, 60 grains.—*London*
Alum, 5 grains, Zinc Sulphate, 5 grains, Basis, 120 grains.—*B P C*

ALUMINIUM ACETATE SOLUTION (*Austr., Belg., Dutch, Ger., and Russ.*)—A clear colourless liquid, with an acid reaction and a faint odour of Acetic Acid, obtained by double decomposition between Aluminium Sulphate and Calcium Acetate. Sp. gr. 1.044 to 1.048. Contains $7\frac{1}{2}$ to 8 p.c. Aluminium Acetate.

This has been incorporated in the *B P C*

Swiss, sp gr 1 055 to 1 059 Contains 10 p c

It is also known as **Burow's Solution**.

A good antiseptic, preferred by some to Carbolic Acid for dressing lacerated wounds

ALUMINIUM ACETO-TARTRATE — Crystals soluble in their own weight of Water Official in Dutch

A powerful, non-poisonous antiseptic, also an astringent caustic

A solution has been sold under the name **Alsol** — *P J* '01, 1 665

A solution containing 10 p c is official in *Swiss*

30 to 60 grains in a pint of Water make a useful gargle or douche.

ALUMINIUM CHLORIDE — A colourless, crystalline mass, giving off fumes of Hydrochloric Acid gas, and becoming damp on exposure to air The crystals form a crystalline powder, chiefly of a yellow colour owing to the presence of iron as an impurity

Dose — 5 to 8 grains = 0 32 to 0 52 gramme

The use of Aluminium Chloride in doses of 5 grains = 0 32 gm. upwards several times a day has proved remarkably efficacious in ataxia — *L* '99, 11 1826

Hydrated Aluminium Chloride is official in *Russ*

Under the names of **Chloralum** and **Chloralum Powder**, preparations containing Aluminium Chloride have been introduced as disinfectants

ALUMINIUM CHLORIDE SOLUTION — Obtained by dissolving Aluminium Hydrate in Hydrochloric Acid A pale yellow liquid. Sp. gr. 1 250 Gargle 12 minims to 1 oz of Water, Spray, 3 minims to 1 oz, Paint, 15 minims to 1 oz Astringent and antiseptic

ALUMINIUM NITRATE — A solution (4 or 6 grains in 1 oz of Water) has been used with success in pruritus vulvæ

ALUMINIUM NAPHTHOL-SULPHONATE (Alumzol) — A whitish powder, readily soluble in Water, introduced as an antiseptic — *P J* (3) xxiii. 605, *C D* '93, 1 94

In treatment of metrorrhagia Used as an intra-uterine injection, with Tincture of Iodine (Alumzol, 2 5, Tincture of Iodine, 25, Absolute Alcohol, 25) by means of an intra-uterine syringe — *M A* '99, 408

Dose — 4 to 8 grains = 0 26 to 0 52 gramme, as an astringent.

ALUMINIUM OLEATE. — A powder Mixed with oil of Lemon, used as a styptic and antiseptic, in checking the progress of eczema — *L* '84, 11 123

ALUMINIUM SULPHATE — White crystalline cakes, or in a white powder, having a sweetish and somewhat astringent taste It is soluble 1 in 1 of Water, insoluble in Alcohol (90 p c) Astringent and antiseptic.

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr., Ger, Jap, Russ, Swed, Swiss and U S

Tests — Aluminium Sulphate dissolves readily in Water, forming a solution which has a strong acid reaction towards blue Litmus paper, this solution yields with Ammonia Solution a white precipitate practically insoluble in excess of the reagent, with Potassium Hydroxide Solution it yields a white gelatinous precipitate soluble in excess of the reagent, but which is again precipitated on the addition of Ammonium Chloride The solution yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid It should contain about 99 p c of pure crystallised Aluminium Sulphate, and shall lose not more than 45 7 p c of its weight when deprived of its Water of crystallisation

It should yield no odour of Ammonia when boiled with Potassium Hydroxide Solution, indicating the absence of Ammonium salts When slightly acidified with diluted Hydrochloric Acid the aqueous solution should not afford an appreciable darkening in colour or a turbidity, on the addition of Hydrogen Sulphide, indicating the absence of more than the slightest traces of Copper

Lead, nor should it be materially darkened in colour on the subsequent addition of Ammonia Solution in slight excess, indicating the absence of more than a trace of Iron

A clear filtered aqueous 10 p c w/v solution of the salt should not become more than faintly opalescent within 5 minutes upon the addition of an equal volume of Tenth-normal Volumetric Sodium Thiosulphate Solution, indicating the limit of free acid

Aluminium Caseinate—A yellowish white, tasteless powder Insoluble in Water Introduced as an intestinal astringent

Dose—5 grains = 0.32 gramme

Salumin Insoluble (Aluminium Salicylate), **Salumin Soluble** (Aluminium and Ammonium Salicylate), **Alkasal** (Aluminium Potassium Salicylate), **Boral** (Aluminium Borotartarate), **Cutol** (Aluminium Borotannate), are preparations containing Aluminium, which have been noticed in medical literature

CIMOLITE—The mineral has yielded on analysis Alumina, 23, Silica, 63, Ferric Oxide, 1.25, Water, 12

A proprietary Toilet article, which is a silicate in very fine powder, is sold under the name 'Cimolite'

FULLER'S EARTH has yielded on analysis Alumina, 10, Silica, 53, Lime, 0.5, Magnesia, 1.25, Ferric Oxide, 9.5, Water, 24

SOAPSTONE, CRETA GALLICA, is a Silicate of Aluminium and Magnesium Is used in prungeo and as a dusting powder for infants, alone or mixed with equal parts of Zinc Oxide or Calamine

AMMONIACUM.

AMMONIACUM

FR, GOMME AMMONIAQUE, GER, AMMONIAGUMMI, ITAL, GOMMA AMMONIACO, SPAN, GOMA AMONIACO

A gum-resin, obtained from *Dorema Ammoniacum*, D Don, and probably other species

It is collected in Persia

Solubility—Sparingly in Water, but forms with it a nearly white emulsion, when 50 grains were digested in 2 oz of Alcohol (90 p c), 40 grains were dissolved, with Alcohol (60 p c) 30 grains were dissolved

Medicinal Properties.—Antispasmodic, stimulant, expectorant, useful in chronic bronchitis and asthma of old people, either in mixture or in pill, as a plaster to promote absorption in chronic synovitis and glandular swellings

Dose.—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—Generally given as *Mistura Ammoniaci*, may be combined with *Tincture of Squill*, or *Fetid Spirit of Ammonia*

Official Preparations—*Emplastrum Ammoniaci cum Hydrargyro* and *Mistura Ammoniaci* Contained also in *Emplastrum Galbani*, in *Pilula Scillæ Composita*, and *Pilula Ipecacuanhæ cum Scillâ*

Not Official—*Pilula Ammoniaci Opiatæ*, *Emplastrum Gummi Resinosum*

Foreign Pharmacopœias.—Official in Austr, Belg, Dan, Dutch, Ger, Hung, Ital, Jap, Mex (*Goma resina Ammoniaco*), Norw, Port, Russ, Span., Swed, and Swiss, Fr., purified by 60 p c Alcohol

Descriptive Notes—Commercial Ammoniacum is imported from Persia, and is believed to be obtained also from *D. Aucheri*, Boiss. It occurs generally in the form of rounded nodules or tears varying in size up to 1 inch (25 mm) in diameter, although averaging only about $\frac{1}{2}$ inch (12 mm). When recently collected the tears are yellowish-white or nearly white, dull externally and opaque internally with a white, slightly polished, fracture, and have an acid, slightly bitter, characteristic taste. Occasionally masses consisting of tears welded together are imported, but these usually contain more or less impurity. When the drug has been long kept the tears assume a brownish-yellow tint.

The official description allows the use of both pale yellow and brownish tears, which may be either white or brownish yellow internally, and they may vary in size from $\frac{1}{4}$ to 1 inch (6 to 25 mm). The odour must not be alliaceous.

Tests—The tests for Ammoniacum are its physical properties, of a Salicylic Acid reaction with Ferric Chloride Solution, a yellow to a brown coloration with Potassium Hydroxide Solution, and an orange-red coloration with Chlorinated Soda Solution.

BP has not yet adopted the determination of the Acid and Saponification values as a means of detecting gums and gum-resins. Notwithstanding the difficulty experienced in sampling and the wide variations between the figures yielded by different specimens, a determination of these constants may often afford a valuable criterion of the purity of a sample. Good commercial Ammoniacum has an Acid value of 92 to 105, and a Saponification value of 145 to 162 according to Dieterich. A sample of good commercial 'tear' Ammoniacum examined in the author's laboratory had an ash limit of 2.15, and gave an Acid value of 106.7, a Resin value of 153.03, a Gum value of 23.87, and a Saponification value of 176.9. A sample of 'mass' Ammoniacum showing an ash limit of 2.55 p.c., gave an Acid value of 101.04, a Resin value of 144.61, a Gum value of 21.06, and a Saponification value of 165.67; 2 samples of powdered Ammoniacum yielding respectively 1.3 and 7.05 p.c. of ash, gave in each instance Acid values of 101.01, Resin values respectively of 150.22 and 153.03, Gum values of 18.26 and 9.83, and Saponification values of 168.48 and 162.86. Dieterich gives the Resin value of commercial varieties as 99.1 to 155.4, the Gum value as 7 to 46.2, and ash of not over 10 p.c.

The more generally occurring impurities are excess of mineral matter and an excessive proportion of matter insoluble in Alcohol (90 p.c.). The absence of Umbelliferone serves to distinguish it from Asafoetida and Galbanum. The *BP* method of performing this test by heating the gum-resin strongly in a dry test tube, cooling, boiling with Water, diluting the resulting liquid largely with Water and making alkaline with Ammonia, is considered unsatisfactory, the result of the test depending greatly on the manner in which the gum-resin is heated. The *PG* test is a more scientific one and is

capable of detecting 2 p c of Galbanum with certainty. The Ammoniacum is boiled with three times its weight of strong Hydrochloric Acid for a quarter of an hour, whereby the Umbelliferone is split off from its natural Ester, the fluid is filtered, and the filtrate supersaturated with Ammonia Solution. It should not exhibit a blue fluorescence when examined by reflected light.

The *B P* gives no indication of the limit of matter insoluble in Alcohol (90 p c), nor of the amount of mineral matter. *P G* on the other hand specifies that the insoluble matter remaining after complete exhaustion of the gum-resin with boiling Alcohol (90 p c) shall amount at the highest to 40 p c, and fixes the ash limit at not more than 5 p c, which is somewhat low. A limit of 7.5 p c of ash has been suggested. The average of a number of good commercial samples examined in the author's laboratory was 6 p c.

Preparations

EMPLASTRUM AMMONIACI CUM HYDRARGYRO. *See* HYDRARGYRUM

As the value of this preparation depends chiefly upon the Mercury it contains, the formula is given under Hydrargyrum.

MISTURA AMMONIACI. AMMONIACUM MIXTURE

Ammoniacum, in coarse powder, $\frac{1}{4}$ oz, Syrup of Tolu, 4 fl drms,
Distilled Water, $7\frac{1}{2}$ fl oz (1 in 32)

Dose — $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc

Foreign Pharmacopœias — Official in Spain (Emulsion), 1 in 33 with White Wine. Not in the others.

Not Official

PILULÆ AMMONIACI OPIATÆ (*Swed*) — Ammoniacum, 10, Myrrh, 5, Squills, 2, Opium, 1, Water, *q s* to make 100 pills.

EMPLASTRUM GUMMI RESINOSUM — Is official in *Dan* and *Norw*. *Swiss* containing 6 p c of Ammoniacum, *Port* (Emplastio Gummo resinoso) containing 2 p c of Ammoniacum. Made with Emplastrum Plumbi.

AMMONIÆ LIQUOR FORTIS.

STRONG SOLUTION OF AMMONIA

A transparent, colourless, very alkaline liquid, with an overpowering pungent smell, containing 32.5 p c (or more correctly 32.35 p c) by weight of Ammonia, NH_3 , eq 16.94.

It should be preserved in well-stoppered glass bottles, which should be kept in a cool atmosphere. Great care should always be exercised in dealing with the liquid.

It may be prepared by the decomposition of an Ammonium salt, usually the Chloride, with Calcium Hydroxide, the resulting gas being dissolved in Water.

In commerce Liquor Ammoniae Fortis is generally sold of sp gr 0.880

K 2

Medicinal Properties.—Usually given in the more diluted form of Solution of Ammonia See below

Official Preparations—Of **Liquor Ammoniae Fortis**, Liquor Ammoniae, Spiritus Ammoniae Fetidus Contained in Linimentum Camphorae Ammoniatum, Linimentum Hydrargyri, and Tinctura Guaiaci Ammoniatum Used in the preparation of Ammonii Benzoas, Ammonii Bromidum, Ammonii Phosphas, Spiritus Ammoniae Aromaticus, and Spiritus Ammoniae Fetidus Of the **Liquor Ammoniae**, Linimentum Ammoniae Contained in Tinctura Ergotae Ammoniatum, Tinctura Opii Ammoniatum, Tinctura Quinine Ammoniatum, Tinctura Valerianae Ammoniatum Used in the preparation of Liquor Bismuthi et Ammonii Citratis, and the scale preparations of Ison

Not Official—Alcohol Ammonia, Lotio Crinalis, Oleate of Ammonia, Liquor Ammoniae Detergens, and Tinct Ammon Comp (Eau de Luce)

Antidotes.—Acetic Acid or Vinegar well diluted with Water, demulcent drinks

Foreign Pharmacopoeias—Official in Belg (Ammonium Hydricum Solutum), sp gr 0.935, 17 p.c., Fr (Ammoniaque Officinale), sp. gr. 0.925, 'U. S. P.' sp gr 0.925, 20 p.c., Mex (Ammoniac), sp gr 0.920, 'U. S. P.' (liquida), sp gr 0.916, Spain (Ammoniac), sp gr. 0.923, U.S. (Aqua Ammoniae Fortior), sp gr 0.897 at 25° C (77° F), 28 p.c., see also Liquor Ammoniae

Tests.—The distinguishing tests are the powerful ammoniacal odour combined with the specific gravity and the volumetric determination figure The *BP* specific gravity is 0.891 at 15.5° C (60° F), the *USP* is 0.897 at 25° C (77° F) It is officially required to indicate 32.35 p.c. by weight of Ammonia (NH_3) as ascertained by titration with Normal Volumetric Solution of Sulphuric Acid It is officially described as containing 32.5 p.c. by weight of NH_3 The *BP* does not mention an indicator; the *USP* gives the choice of Litmus or Methyl Orange Solution

So far as impurities are concerned Liquor Ammoniae Fortis should respond to the tests given under Liquor Ammoniae, *qv*

Aqua Ammoniae Fortior *USP* when diluted with twice its volume of Distilled Water should respond to the *USP* volumetric test given under Liquor Ammoniae The *PG* recognises only the dilute Ammonia

LIQUOR AMMONIAE. SOLUTION OF AMMONIA.

A clear, colourless liquid, containing 10 p.c. by weight of Ammonia, NH_3 ; prepared by mixing 1 of strong Solution of Ammonia with 2 of Distilled Water

It possesses a characteristic pungent odour, a powerfully caustic taste, and strongly alkaline reaction

It should be preserved in well-stoppered bottles, which should be kept in a cool atmosphere

Medicinal Properties—A general stimulant Externally (applied to the nostrils) in syncope, an excellent application to the sting of a wasp or the bites of insects On the skin it is a powerful rubefacient, and in embrocations it is used as a counter-irritant for pain, stiffness of joints, bronchitis, etc Was at one time used by injection as an antidote to snake bites, but Potassium Permanganate is now considered a better antidote.

Dose —10 to 20 minims = 0·6 to 1·2 c c, well diluted

Prescribing Note —*Ammonia is more generally prescribed internally in the form of Spiritus Ammonia Aromaticus or of Ammonium Carbonate*

Official Preparations —*Linimentum Ammonia* Used in the preparation of *Ammonii Benzoas*, *Ferri et Ammonii Citras*, *Ferri et Quininae Citras*, *Ferrum Tartaratum*, *Liquor Bismuthi et Ammonii Citratis*, *Tinctura Opii Ammoniata*, *Tinctura Quininae Ammoniata*

Not Official —*Liquor Ammonii Anisatus*, *Spiritus Ammonii Anisatus*, *Spiritus Ammonia Foeniculi* *utis*, *Spiritus Ammonia*

Foreign Pharmacopœias —Official in Austr., Dan., Dutch, Fr. (*Ammoniaque Diluée*), Ger., Hung., Jap., Norw., Russ., Swed., Swiss, 10 p c, U.S. (*Aqua Ammonia*), 10 p c, sp gr 0·958 at 25° C (77° F), Belg., Fl., Ital., Mex., Port., Span. and U.S., see *Ammon. Liq. Fort.*

Tests —The distinguishing tests for *Liquor Ammonia* are the pungent ammoniacal odour, the sp gr at 15·5° C (60° F), which should be 0·959 [0·958 at 25° C (77° F)], *USP*, 0·960, *PG*], and that it is officially required to indicate 10 p c by weight of Ammonia (NH_3) when titrated with Normal Volumetric Sulphuric Acid Solution. Neither *BP* nor *PG* make any reference to the indicator of neutrality to be used, the *USP* states Litmus or Methyl Orange Test-solution, the latter is usually employed, Phenolphthalein Solution being useless.

The *BP* volumetric test indicates 10·0 p c by weight of Ammonia (NH_3), the *PG* 9·95 p c to 10·0 p c by weight. *USP* defines it as containing 10 p c by weight of Ammonia gas.

When a glass rod moistened with Hydrochloric Acid is brought near Ammonia Solution dense white fumes of Ammonium Chloride are given off. Ammonia Solution when sufficiently highly diluted yields on the addition of Potassio-mercuric Iodide (Nessler's) Solution a characteristic brown coloration, which in the presence of much Ammonia changes to a brown or reddish-brown precipitate.

The more generally occurring impurities are empyreumatic and mineral matter, heavy metals, *eg*, Arsenic, Iron, Lead, and Zinc, Ammonium Carbonate or Carbamate, Calcium Carbonates, Chlorides, and Sulphates. The *BP* specifically mentions a test for absence of tarry matters, requiring that no colour or odour should be produced on the addition of a slight excess of Hydrochloric Acid to a mixture of equal volumes of Ammonia and Water, but is content to group the remainder without any regard to their relative importance under the expression 'it shall yield no characteristic reaction for,' etc. When rendered faintly acid by the addition of Hydrochloric Acid it should be unaffected by Hydrogen Sulphide Solution, indicating the absence of Arsenic and Lead. On subsequently rendering the solution again alkaline by the addition of Ammonia Solution no perceptible darkening in colour or turbidity should be produced, indicating the absence of more than a faint trace of Iron or Zinc. The solution should not effervesce on the addition of diluted Hydrochloric Acid, indicating the absence of Carbonates, when almost neutralised with Hydrochloric Acid it should not yield an opalescence on the addition of Ammonium Oxalate Solution, indicating the absence of Calcium. When super-saturated with Nitric Acid it shall yield little or no turbidity on the

addition of Silver Nitrate Solution, nor on the addition of Barium Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. The *USP* and *PG* require the neutralised acid to yield, on evaporation, a residue which is completely volatilised on ignition, the *PG* also requires the residue to be colourless. No reference to fixed or volatile character of the residue left on evaporation occurs in the *BP*. The *PG* requires that when diluted with 4 times its volume of Calcium Hydroxide Solution it should show only a faint turbidity after the lapse of one hour, indicating the absence of Ammonium Carbonate and Carbamate. No test similar to this appears in the *BP*. The test with Decinormal Volumetric Potassium Permanganate Solution for readily oxidisable organic impurities is peculiar to the *USP*. If 0.1 c.c. of Tenth-normal Volumetric Potassium Permanganate Solution be added to 10 c.c. of Ammonia Solution, slightly supersaturated with diluted Sulphuric Acid, the pink colour should not be completely destroyed within 10 minutes.

Volumetric Determination.—1 gramme neutralises 59 cc of VS of Sulphuric Acid, *BP*, 5 cc should require 28 to 28.2 cc Normal VS. of Hydrochloric Acid, *PG*, the *USP* gives the following directions for making the determination —Introduce into a stoppered weighing bottle 3 cc of Ammonia Water and weigh accurately Dilute with 50 cc of Distilled Water and titrate with normal VS of Sulphuric Acid, using Litmus or Methyl Orange TS as indicator Multiply the number of cc of the VS of Sulphuric Acid consumed by 1.693, and divide this product by the weight of the Ammonia Water taken, the quotient represents the percentage of Ammonia gas

Preparations.

LINIMENTUM AMMONIÆ. LINIMENT OF AMMONIA.

Solution of Ammonia, 1, Almond Oil, 1, Olive Oil, 2	Mix by
shaking	(1 in 4)

Cotton Seed, Sesame and Nut Oils have each been recommended, but Cotton Seed is the only Oil which makes a satisfactory and permanent Emulsion

Foreign Pharmacopœias.—Official in Austr, Dutch, Hung and Jap, 1 and 4 Sesame Oil, Belg, 1 and 9 Medicinal Oil, Fr, 1 and 9 Olive Oil, Ger, Liq Am 1, Olive Oil 3, Poppy Oil 1, Ital, 1 and 4 Olive Oil, Mex, 1 Sesame Oil 9, also 1, sesame Oil 4; Port, 1 and 4 Almond Oil, Russ, Liq. Am 1, Olive Oil 3 Sesame Oil 1, Span, 1 and 9 Olive Oil, Swed, 1 and 3 Olive Oil, Swiss, 1 and 3 Sesame Oil, U S, Am 35, Alcohol 5, Cotton Seed Oil 57, Oleic Acid, 3. Not in Norw. All by weight, except U S.

SPIRITUS AMMONIÆ AROMATICUS. See AMMONII CARBONAS.

SPIRITUS AMMONIÆ FETIDUS FETID SPIRIT OF AMMONIA

Asafetida, $1\frac{1}{2}$, strong Solution of Ammonia, 2, Alcohol (90 p c),
to make 20. (14 in 20)

Nervine stimulant and antispasmodic, useful in hysteria

Dose.—For repeated administration, 20 to 40 minims = 1·2 to 2·4 c c , for a single administration, 60 to 90 minims = 3·6 to 5·3 c c .

Tests.—A clear, almost colourless liquid, possessing a pungent

ammoniacal and alliaceous odour. It has a specific gravity of 0.848, and it is officially required to contain 2.88 grammes of Ammonia (NH_3) per 100 c.c. as indicated by titration with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as an indicator of neutrality. The *B.P.* does not include a figure for specific gravity, but gives the volumetric test, it, however, does not state what indicator of neutrality is to be employed.

Not Official

ALCOHOL AMMONIA—Absolute Alcohol saturated with Ammonia gas. It contains about 14 p.c. of NH_3 .

It is used in filling and renovating Smelling Salt bottles.

LOTIO CRINALIS—Ol. Amygdal., 1 fl. oz., Liq. Ammon. Fort., 1 fl. oz., Sp. Rosmar., 4 fl. oz., Aq. Mellis, 2 fl. oz.

This has been incorporated in the *B.P.C.* as follows—

Almond Oil, 12.50, Strong Solution of Ammonia, 12.50, Oil of Rosemary, 0.50, Alcohol, 50, Honey Water, *q.s.* to produce 100.

SPIRITUS or LIQUOR AMMONII ANISATUS

Austr. and *Ger.*—Anethol, 1, Alcohol, 24, Solution of Ammonia, 5.

Belg. (Ammonia Spiritus Anisatus)—Anethol, 8, Alcohol, 77, Solution of Ammonia, 20.

Hung. and *Russ.*—Oil of Anise, 1, Alcohol, 24, Solution of Ammonia, 6.

Ital. and *Span.*—Oil of Anise, 1, Alcohol, 24, Solution of Ammonia, 5.

Dan., *Norw.* and *Swed.*—Oil of Anise, 1, Alcohol, 32, Solution of Ammonia, 7.

Dutch.—Oil of Anise, 1, Alcohol, 19, Solution of Ammonia, 5.

Swiss.—Oil of Anise, 3, Alcohol, 77, Solution of Ammonia, 20.

All by weight.

B.P.C. (Liquor Ammonia Anisatus)—Anethol, 3.50, Solution of Ammonia, 16.50, Alcohol, *q.s.* to produce 100.

Spiritus Ammonia Foeniculatus—Oil of Fennel, 3, Alcohol, 80, Ammonia Water, 17—*Jap.*

Spiritus Ammonia—Stronger Ammonia Water, 250, Alcohol, *q.s.* to make the product contain 10 p.c. by weight of Ammonia Gas—*U.S.*

OLEATE OF AMMONIA—Oleic Acid, 1 oz., Spirit, 1 oz., Solution of Ammonia, 7 oz., Distilled Water, to 16 oz. Pour the acid into a bottle, mix the Spirit and Ammonia, and pour into the bottle. Cork tightly, and allow to stand a week or more until saponification is complete. This is suitable for adding to Solution of Ammonia (1 to 8) to make a household article—*Pharm. Form.*

Liquor Ammonia Detergens *Syn.* Household Ammonia—Strong Solution of Ammonia, 30, Oleic Acid, 6, Alcohol, 6, Distilled Water, *q.s.* to produce 100—*B.P.C.*

Note—If a 'cloudy' preparation be desired, about half of the Distilled Water in the above formula should be replaced by hard tap Water, the exact proportion depending upon the amount of total solids in the hard Water—*B.P.C.*

TINCTURA AMMONIAE COMPOSITA EAU DE LUCE—Mastic, 2 drms., Alcohol (90 p.c.), 9 fl. drms., Ol. Lavand., 14 minims, Liquor Ammonia Fortis, 20 fl. oz.

This has been incorporated in the *B.P.C.* as follows—

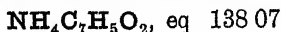
Mastic, 1.25, Alcohol, 5.50, Oil of Lavender, 0.15, Strong Solution of Ammonia, *q.s.* to produce 100.

Stimulant, antispasmodic. Has been used in tropical countries as an application to snake bites.

Dose.—5 to 10 minims = 0.3 to 0.6 c.c., in Water.

AMMONII BENZOAS.**AMMONIUM BENZOATE**

FR, BENZOATE D'AMMONIUM, GER, AMMONIUMBENZOAT, ITAL, BENZOATO DI AMMONIO, SPAN, BENZOATO AMONICO



Fine, white laminar crystals, or a crystalline powder, odourless or possessing a faint odour of Benzoin, and a saline taste. It is produced by the combination of Benzoic Acid with Ammonia. It gradually loses Ammonia on exposure to air.

Solubility—1 in 6 of Water, 1 in 22 of Alcohol (90 p.c.), 1 in 8 of Glycerin.

It will not quite dissolve 1 in 5 of Water, as sometimes stated.

Medicinal Properties.—Valuable in chronic vesical catarrh with alkaline urine and phosphatic deposit, and in chronic bronchial catarrh with much secretion. It is more soluble than Benzoic Acid, and therefore should be preferred, and is less irritant to the alimentary canal.

An intestinal antiseptic in typhoid—*M A '94*, 555

Dose.—5 to 15 grains = 0.32 to 1 gramme

Prescribing Note.—*Usually given in solution*

Incompatibles.—Acids, Liquor Potassæ, and Ferric salts

Foreign Pharmacopœias—Official in Jap, Mex, Port, Russ, Span, Swiss and U.S.

Tests.—The distinguishing tests for Ammonium Benzoate are the odour of Ammonia, which is evolved when its aqueous solution is heated with Solution of Potassium or Sodium Hydroxide, the yellowish-brown coloration, produced when its sufficiently diluted solution is mixed with Potassio-mercuric Iodide (Nessler's) Solution, and the buff-coloured precipitate thrown down when its sufficiently concentrated aqueous solutions are acidified with a mineral acid, and the characteristic buff-coloured precipitate produced on adding Test-solution of Ferric Chloride to its aqueous solution. The *U.S.P.* requires that it should contain not less than 98 p.c. of pure Ammonium Benzoate, but does not state a method of determination.

The more generally occurring impurities are mineral matter, shown by a residue being left on ignition, free Benzoic Acid, indicating imperfections in the process of manufacture, and detected by the reaction of its cold aqueous solution towards blue Litmus paper or Solution, it should be neutral or only slightly acid, Chlorides and Sulphates, which indicate Toluene or Hippuric Acid as its probable source, and which are detected by the usual tests for Chlorides and Sulphates after removal of the Benzoic Acid. The 10 p.c. w/v aqueous solution is acidified with diluted Nitric Acid, and the precipitated Benzoic Acid is separated by filtration, the filtrate should be unaffected by the addition of either Silver Nitrate or Barium Chloride Solution. The *B.P.* includes tests for all these impurities. *U.S.P.* includes heavy metals as a likely impurity, and requires that the

acidulated, filtered 5 p c aqueous solution of the salt shall respond to the time-limit test for heavy metals. It also requires that the Benzoic Acid prepared from the salt shall answer the tests and be free from the impurities given under Benzoic Acid. No such requirement appears in the *BP*, either under this heading or under Sodium Benzoate.

Residue—On strongly heating it emits vapours having the odour of Ammonia and Benzoic Acid, and is finally volatilised, *USP*, at a red heat it leaves no residue, *BP*.

Not Official

AMMONII BORAS

A crystalline salt, with an alkaline reaction

Solubility—1 in 15 of Water

Medicinal Properties—Has been used with success in renal and vesical calculi

For renal colic, 20 grains = 1.3 grammes, every two hours until free passage of urine takes place, then 15 grains = 1 gramme, three times a day—*TG* '87, 623

AMMONII BROMIDUM.

AMMONIUM BROMIDE

NH_4Br , eq 97.29

Fr, Bromure d'Ammonium, Gr, Ammoniumbromid, Ital, Bromuro di Ammonio, Span, Bromuro Amonico

Small, colourless and odourless, prismatic crystals, or a white crystalline powder possessing a pungent saline taste

It is prepared by the neutralisation of Hydrobromic Acid by Ammonia

Solubility.—1 in $1\frac{1}{2}$ of Water, and measures 2, 1 in 15 of Alcohol (90 p c)

Medicinal Properties—An excellent nervine sedative and depressant, especially useful for sleeplessness, the result of worry or mental anxiety and fatigue, anaphrodisiac, given in epilepsy, with Chloral in acute alcoholism, in acute mania and nymphomania, and in many other conditions in which the Potassium salt is used. Not so apt to produce Bromism as the Potassium salt, and less depressing. Relieves headache, especially in migraine, and neuralgic pain. Sedative in pharyngeal and laryngeal irritation. Useful in whooping-cough and asthma.

Reference to the use of this salt in the treatment of epilepsy appears in *L* '05, 1710. If benefit does not follow a daily dose of from 45 to 60 grains of one or a combination of the Bromide salts in epilepsy, some other remedy or method of treatment should be sought. In confirmed epilepsy with mental deterioration, all that can be expected from the continuous use of the Bromides is diminution in the number and perhaps in the severity of the seizures. In serial epilepsy and the status epilepticus Chloral in combination with the Bromides forms the most effective remedy.

Dose.—5 to 30 grains = 0.32 to 2 grammes

Incompatible—spirit of Nitrous Ether

Not Official Preparations—Ammonium Bromide Effervescent, Elixir Ammonii Bromidi, Pastilles Ammonii Bromidi, Trochisci Ammonii Bromidi

Foreign Pharmacopœias—Official in Austr., Dan., Dutch, Fr., Ger., Ital., Jap., Mex (Bromuro de Amonio), Norw., Russ., Span., Swed., Swiss and U.S. Not in the others

Tests.—The following tests for Ammonium Bromide are that an aqueous solution of the salt shall, when heated with Sodium or Potassium Hydroxide Solution, evolve Ammonia, the latter is readily recognised by its odour or its immediately turning a piece of moistened red Litmus paper blue, the sufficiently diluted aqueous solution yields a yellowish-brown coloration with Potassio-mercuric Iodide (Nessler's) Solution. The aqueous solution of the salt should yield with Silver Nitrate Solution a yellowish-white precipitate insoluble in Nitric Acid, practically insoluble in dilute Ammonia Solution, but soluble in Potassium Cyanide Solution. On the addition of Chlorine Solution to an aqueous solution of the salt, Bromine is set free, which on shaking with a little Chloroform or Carbon Bisulphide yields a reddish solution. It is officially required to contain not less than 99.43 p.c. nor more than 100.79 p.c. of pure Ammonium Bromide, 0.5 gramme of the salt requiring, when dissolved in Water, not less than 51.1 c.c. nor more than 51.8 c.c. of Volumetric Silver Nitrate Solution. The *U.S.P.* requires it to contain not less than 97 p.c. of Ammonium Bromide, the *P.G.* not more than 100.98 p.c. The volumetric determination adopted by the *B.P.* is carried out on the dry salt, but no indication of the limit of Water permissible is given.

The *P.G.* also employs a salt dried at 212° F (100° C) in carrying out the volumetric determination. The *U.S.P.* does not direct the salt to be dried.

The more generally occurring impurities are heavy metals, *e.g.*, Copper, Lead and Iron, Barium, Bromates, Chlorides, Iodides and Sulphates. A 5 p.c. aqueous solution of the salt when slightly acidified with Hydrochloric Acid should be unaffected by Hydrogen Sulphide Solution, indicating the absence of Copper and Lead. The *U.S.P.* and the *P.G.* include a separate test for Iron with Potassium Ferrocyanide Solution, which is given in the small type below. Barium, Bromates, Iodides and Sulphates may, if present, be detected by the tests given under the respective headings of Potassium Sulphate Solution, Diluted Sulphuric Acid, Chlorine Water, Chloroform and Barium Nitrate Solution. The precipitate produced on the addition of Silver Nitrate Solution to an aqueous solution of the salt should be practically insoluble in Ammonia, and the filtered ammoniacal liquid should yield little or no opalescence when supersaturated with Nitric Acid, indicating the absence of more than a trace of Chlorides. The salt should be entirely volatilised on heating leaving a weighable residue, indicating the absence of fixed matter.

Chlorine Water and Chloroform.—If Chloroform be added to an aqueous solution of the salt (1 c.c. Chloroform and 10 c.c. of a (1-20) solution, *U.S.P.* and Chlorine Water (diluted with an equal volume of Water, *U.S.P.*) be

carefully introduced with constant agitation, the Chloroform is coloured reddish brown, *P G*, yellow to orange, *U S P*, free from any violet tint, *U S P*

Diluted Sulphuric Acid.—A small quantity of the salt spread out on a porcelain slab should not at once assume a yellow colour on the addition of a few drops of diluted Sulphuric Acid (test for Bromate), *P G* and *U S P*

The aqueous solution (1 in 20) should be unaffected by diluted Sulphuric Acid, *P G*

Barium Nitrate Solution—The aqueous solution should be unaffected by Barium Nitrate Solution, *P G*

Potassium Sulphate Solution—There should be no turbidity in 10 c c of a (1-20) aqueous solution acidulated with Acetic Acid, on the addition of 1 c c Potassium Sulphate Solution, *U S P*

Potassium Ferrocyanide Solution—20 c c of an aqueous solution (1-20) should not be immediately turned blue on the addition of 0.5 c c Potassium Ferrocyanide Solution, *P G* *U S P* uses a 1-100 solution of the salt, but does not state test quantities

Volumetric Determination—10 c c of a solution of 3 grammes in 100 c c of Water, with the addition of a few drops Potassium Chromate Solution (1 drop, *P G*) should require not more than 30.9 c c (*P G*), 31.6 c c (*U S P*), of the Deci normal Volumetric Solution of Silver Nitrate to produce a permanent red colour, the *U S P* directs the salt itself to be used for titration without drying, the *P G* uses the salt previously dried at 100° C (212° F)

Not Official

AMMONII BROMIDUM EFFERVESCENS, is made of 2 strengths containing 5 and 10 grains in 60 grains

ELIXIR AMMONII BROMIDI—Ammonium Bromide, 85, Citric Acid, 4, Aromatic Elixir (*U S P*), *q s* to make 1000—*U S N F* 1896

In *U S N F* 1906 the formula remains the same without the Citric Acid, which is omitted

Contains 5 grains of Ammonium Bromide in each fl dr m
Ammonium Bromide, 10, Citric Acid, 0.50, Aromatic Elixir, *q s* to produce 100—*B P C*.

Contains about 5½ grains in each fl dr m

The *B P C* appears to contain twice as much Compound Spirit of Orange as does the *U S P*, but as the *B P C* Compound Spirit is half the strength, the flavouring of the two Elixirs is about the same. The alterations in the *B P C Supplement* leave the result much the same

LOZENGES, containing 2 grains=0.13 gramme, of Ammonium Bromide in each. Useful in whooping cough

Dose—1 to 3 lozenges

Pastilles containing 1 grain in each with Glyco gelatin Basis

AMMONII CARBONAS.

AMMONIUM CARBONATE

FR, CARBONATE (SESQUI) D'AMMONIAQUE, GER, AMMONIUMCARBONAT, ITAL, CARBONATO DI AMMONIO, SPAN, CARBONATO AMONICO

A mixture of Ammonium Hydrogen Carbonate, NH_4HCO_3 , with Ammonium Carbamate, $\text{NH}_4\text{NH}_2\text{CO}_2$

Hard, transparent, crystalline masses, possessing a strong ammoniacal, but not empyreumatic, odour and strong alkaline reaction. It effloresces when exposed to the air, and becomes covered on the surface with a white powder. On this account it should be kept in

well-stoppered bottles and in a cool atmosphere Only the translucent portions of the mass should be used for dispensing purposes

Solubility.—1 in 4 of Water; 1 in 200 of Alcohol (90 p c), 1 in 5 of Glycerin

Medicinal Properties.—Gastric, cardiac, and general stimulant, a valuable expectorant, frequently combined with Ipecacuanha in acute and chronic bronchitis when the phlegm is tough and scanty Rarely as an emetic in $\frac{1}{2}$ drm doses

Has been recommended in full and continuous doses in cholera, in the place of alcoholic stimulants

Dose.—3 to 10 grains = 0.2 to 0.65 gramme

Prescribing Note.—15 grains dissolved in Water are taken with 17 grains of Citric Acid to form a saline draught

Incompatibles.—Acids, Acid salts, Iron salts, Lime Water, and salts of the alkaline earths and of the alkaloids

Official Preparations.—Used in the preparation of Ammonii Chloridum, Bismuthi Carbonas, Ferri Carbonas Saccharatus, Liquor Ammonii Acetatis, Liquor Ammonii Citratis, and Spiritus Ammoniae Aromaticus

Not Official.—Linctus Ammoniae Compositus, Linctus Ammonii Acetatis Fortior, Liquor Ammonii Citratis Fortior, Liquor Volcani Cervi or Spiritus of Hartshorn, Mistura Ammoniae cum Senega, Hartshorn and Oil, and Ammonium Bicarbonate

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Port., Russ., Span., Swiss and U.S.

Tests.—The distinguishing tests for Ammonium Carbonate are its strong ammoniacal odour and alkaline reaction, when heated, either alone or with Potassium or Sodium Hydroxide Solution, it evolves Ammonia, its sufficiently diluted aqueous solution yields a yellowish-brown coloration with Nessler's reagent (Potassio-mercuric Iodide Solution), it effervesces with dilute mineral acids, evolving a gas which causes a white precipitate with Lime Water, and its aqueous solution yields with Barium Chloride Solution a white precipitate, soluble with effervescence in diluted Hydrochloric Acid

Each gramme is officially required to neutralise not less than 18.7 c.c. of Volumetric Sulphuric Acid Solution, corresponding to 97.25 p.c. of a salt of the pharmacopœial composition This figure is considered too high even for the best specimens, and it has been suggested (*P.J.* '01, 1775) that the figure should be altered to 18.0 c.c. The *U.S.P.* requires that it should contain not less than 97 p.c. of a mixture of Acid Ammonium Carbonate and Ammonium Carbamate, and that it should yield not less than 31.58 p.c. of Ammonia gas. The method adopted by the *U.S.P.* for its volumetric determination is given in the small type below

The more generally occurring impurities are empyreumatic and mineral matter, Lead, Copper, and Iron, Chlorides, sulphates and Thiosulphates

The presence of empyreumatic or non-volatile matter is detected by the appearance and odour of the residue left on the evaporation of the neutralised salt, and by any residue left upon gentle ignition. The test with Silver Nitrate Solution for Thiosulphate and limit of

Chloride is peculiar to the *P G* and *U S P*. A 1 in 20 aqueous solution of the salt when rendered faintly acid by the addition of Hydrochloric Acid should not be affected by Hydrogen Sulphide Solution, indicating the absence of Lead and Copper, nor on the subsequent addition of Ammonia Solution should it be darkened in colour, indicating the absence of Iron. A standard of 5 parts per 1,000,000 has been suggested (*C D* '08, 1795) as a standard for Lead, Arsenic not having been found in this chemical. Chlorides, Sulphates and Thio-sulphates may be detected, if present, by the tests with Silver Nitrate Solution and Barium Nitrate or Chloride Solution given in the small type below. The *P G* includes a test for Calcium, and requires that a 1 in 20 aqueous solution should be unaffected by Ammonium Oxalate Solution. It also gives a test with Ferric Chloride Solution, requiring that a 1 in 20 aqueous solution of the salt should not be coloured red on the addition of Ferric Chloride Test-solution.

Residue—On heating, Ammonium Carbonate is volatilised, and should leave no residue, *B P*, *P G*, and *U S P*. If an aqueous solution (1 gramme of the salt, *P G* and *U S P*) be supersaturated with Nitric Acid, and evaporated to dryness, the residue should be colourless and odourless, *B P*, *P G*, and *U S P*, and on gentle ignition should be completely volatilised, *P G* and *U S P*.

Silver Nitrate Solution—An aqueous solution (1–20) of Ammonium Carbonate should neither assume a brown colour, nor become more than slightly opalescent within two minutes, on the addition of Silver Nitrate Solution and subsequent supersaturation with Nitric Acid, indicating the absence of Thio sulphate and limit of Chloride, *P G* and *U S P*.

Barium Nitrate or Barium Chloride Solution—An aqueous solution (1 in 20) should be unaffected by Barium Nitrate Solution, *P G*, or by Barium Chloride Solution, *U S P*.

Volumetric Determination—2 grammes of the unaltered translucent salt, dissolved in a mixture of 50 c c each of Water and Normal V S Sulphuric Acid, and then boiled for a few minutes to expel the liberated CO_2 , should, when the solution is cooled, require not more than 12.7 c c of Normal V S Potassium Hydroxide for exact neutralisation, Litmus T S being used as indicator, *U S P*.

Preparations

SPIRITUS AMMONIÆ AROMATICUS AROMATIC SPIRIT OF AMMONIA *B P Syn*—SPIRITUS AMMONIÆ COMPOSITUS SPIRIT OF SAL VOLATILE

A clear, almost colourless liquid, possessing a strong ammoniacal odour and taste. It gradually darkens on exposure to light, and on this account should be kept in well-stoppered bottles of a dark amber tint and in a cool atmosphere.

It is prepared by mixing Oil of Nutmeg, $4\frac{1}{2}$ fl drms, Oil of Lemon, $6\frac{1}{2}$ fl drms, Alcohol (90 p c), 120 fl oz, and Distilled Water, 60 fl oz, and distilling until 140 fl oz has been collected. This portion is reserved and a further 9 fl oz is distilled, this second distillate is transferred to a strong, well stoppered bottle, mixed with 8 fl. oz of Strong Ammonia Solution and 4 oz of Ammonium Carbonate, and allowed to stand at 60°C (140°F) until the Ammonium Carbonate is dissolved, when the solution is filtered into the distillate first reserved. The official directions are that 149 fl oz should be distilled, collecting separately and reserving the last 9 fl oz passing

over, but it is more convenient to dissolve the Ammonium Carbonate and Ammonia in 9 fl oz of Water while the distillation is proceeding, and not to carry it past 140 fl oz

Medicinal Properties.—Similar to those mentioned under Ammonium Carbonate. A domestic remedy for nervous headache, more useful when combined with Ammonium Bromide

Dose.—20 to 40 minims = 1.2 to 2.4 c.c., for repeated administration, for a single administration, 60 to 90 minims = 3.6 to 5.3 c.c.

Foreign Pharmacopœias—(*Spiritus Ammonia Aromaticus*) Jap., Ammonium Carbonate 40, Ammonia Water 100, Oil of Lemon 8, Oil of Cloves 1, Oil of Lavender 1, Alcohol 650, Distilled Water 200, U.S., Ammonium Carbonate 84, Ammonia Water 90, Oil of Lemon 10, Oil of Lavender 1, Oil of Cloves 1, Alcohol 700, Distilled Water, *qs* to make 1000. Neither are distilled. Port (*Esprit Ammoniacal Aromatique*), *distilled*, contains Carbonate Austr., Dan., Dutch, Ger., Hung., Ital., Norw., Russ., Span., Swed. and Swiss, have Liquor or Spiritus Ammonii Anisatus, a mixture of Oil of Anise, Spirit, and Liq. Ammon., but in slightly different proportions, Belg., a mixture of Anethol, Spirit and Liq. Ammon. See p. 135

Tests.—The distinguishing tests for Spiritus Ammonia Aromaticus are its strong ammoniacal odour and taste, the specific gravity, which should be between 0.888 and 0.893. The addition of 16 c.c. of Barium Chloride Solution to 20 c.c. of the spirit should yield a precipitate, becoming more copious on heating to 71° C (160° F), and the filtrate from this precipitate should, on the addition of a further quantity of Barium Chloride Solution and again warming, again yield a precipitate.

The above test with Barium Chloride Solution is generally considered unreliable. It has been shown (*P.J.* '00, i 147) that the precipitation of Barium Carbonate in the presence of Ammonium salts by Barium Chloride does not form a satisfactory basis for the determination of Ammonium Carbonate in the aromatic spirit, and the somewhat complicated method of measuring the Carbonic Acid gas in a nitrometer is suggested. The necessity for resorting to this latter method can be, however, obviated, as it has been pointed out (*P.J.* '00, ii 105) that the addition of Ammonium Chloride to the solution alters the character of the precipitate altogether. A weighed quantity of 5 grammes of solid Ammonium Chloride is added to the 20 c.c. of aromatic spirit, and after vigorous agitation the requisite quantity of Barium Chloride Solution is added. The mixture is warmed to 71.1° C (160° F), cooled to the normal temperature and filtered. The filtrate, on the addition of more Barium Chloride Solution and warming, gives no further precipitate.

It is officially required to contain about 2.4 p.c. by weight of Ammonia gas, equivalent to 2.16 grammes in 100 c.c., as ascertained by titration with Volumetric Solution of Sulphuric Acid. The *B.P.* does not state the indicator of neutrality to be used; Methyl Orange Solution is suitable, 20 c.c. of the Spirit neutralises 25 c.c. of the Volumetric Acid.

The official test for total alkalinity is shown (*P.J.* '00, i 145) to make no allowance for deterioration of the spirit during the process

of manufacture or during storage, the latter obviously being a variable quantity, depending upon a number of variable conditions. The addition of a few c c per litre of the strong Solution of Ammonia is suggested in the same reference as a means of sufficiently raising the total alkalinity.

LIQUOR AMMONII ACETATIS. SOLUTION OF AMMONIUM ACETATE

1 of Ammonium Carbonate dissolved in 10 of Distilled Water, neutralised with Acetic Acid, and diluted with Distilled Water to make 20

This dilute solution is now prepared direct from Ammonium Carbonate as recommended in the *Companion* 1894, and the concentrated solution is omitted.

Medicinal Properties—Diaphoretic, diuretic and slightly antipyretic. A mixture of this medicine with Spirit of Nitrous Ether forms one of the oldest remedies for febrile conditions, and, there being no risk of its producing collapse, one of the safest. Given in full doses for alcoholism.

Dose—2 to 6 fl drms = 7 l to 21 3 c c

Incompatibles—Potassium and Sodium Hydroxides, and alkaline Carbonates.

Foreign Pharmacopœias—Official in Port, sp gr 1 029, Fr, sp gr 1 036, Mex and US, all made with Carbonate, Austr, sp gr 1 030, Ital, 1 034, Norw, sp gr 1 035 to 1 040, Belg, Dutch, Ger, Hung, Jap, Russ and Swiss, sp gr 1 032 to 1 034, Span, sp gr 1 036 all made with Caustic Ammonia.

Tests—A clear, almost colourless fluid possessing a faint acetous odour and faint saline, acidulous but not empyreumatic taste. It should have a specific gravity of about 1 018. The *B.P.* states that a small quantity of the liquid, when deprived of its Carbonic Anhydride by heating in a test-tube, shall possess a neutral reaction to test-papers. Solution of Cochineal affords a more useful means of determining the neutrality of the solution, and obviates the necessity of boiling off the Carbonic Anhydride.

LIQUOR AMMONII CITRATIS — SOLUTION OF AMMONIUM CITRATE

Citric Acid 5, dissolved in Distilled Water 25, neutralised with Ammonium Carbonate, and diluted with Distilled Water to make 40

Medicinal Properties—Similar to Liquor Ammonii Acetatis

Dose—2 to 6 fl drms = 7 l to 21 3 c c

Tests—A clear, almost colourless and odourless liquid possessing a saline taste. It should have a specific gravity of about 1 057. The remarks upon the method of determining the neutrality of the solution appearing upon Liquor Ammonii Acetatis apply equally here.

Not Official.

LINCTUS AMMONIÆ COMPOSITUS — Ammonium Carbonate, $\frac{1}{2}$ grain, Ipecacuanha Wine, 2 minims, Tincture of Squill, 5 minims, Essence of Anise, 1 minim, Mucilage of Acacia, 20 minims, Water, to 1 fl. drms — *Royal Chest.*

LIQUOR AMMONII ACETATIS FORTIOR.—Carbonate of Ammonium, 15½ oz, Acetic Acid, 50 fl oz or *q s*, Distilled Water, *q s* to make 60 fl oz.
B P 1885

This has been incorporated in the *B P C* as follows —

Ammonium Carbonate, 25, Acetic Acid, *q s* to neutralise, Distilled Water, *q s* to make 100

LIQUOR AMMONII CITRATIS FORTIOR —Citric Acid, 12 oz, Strong Solution of Ammonia, 11 fl oz or *q s*, Distilled Water *q s* Neutralise the acid with the Ammonia, adding sufficient Distilled Water to make 24 fl oz —*B P* 1885.

This has been incorporated in the *B P C*

LIQUOR VOLATILIS CORNU CERVI, or SPIRIT OF HARTSHORN —Solution of Carbonate of Ammonia of the old Pharmacopœias, distilled from Hartshorn, but is now more generally represented by Liquor Ammoniac
B P

MISTURA AMMONIÆ CUM SENEGA — Ammonium Carbonate, 4 grains, Ipecacuanha Wine, 10 minims, Infusion of Senega, ½ fl oz; Water, to 1 fl oz —*St Thomas's*

This has been incorporated in the *B P C*, using Ammonium Carbonate 5 grains in the place of 4

Ammonium Carbonate, 5 grains, Tincture of Squill, 12 minims, Spirit of Chloroform, 10 minims, Infusion of Senega, to make 1 fl. oz —*Royal Rec*

HARTSHORN AND OIL —1 of Sp Hartshorn and 3 of Oil of Almonds

AMMONIUM BICARBONATE —White, crystalline powder Soluble 1 in 5 of water, insoluble in Alcohol (90 p c) It is formed when Ammonium Carbonate is exposed to the air Employed in powders and pastilles as a substitute for Ammonium Carbonate

AMMONII CHLORIDUM.

AMMONIUM CHLORIDE

N O Syn —AMMONIUM CHLORATUM, CHLORETUM AMMONIUM

FR, CHLORURE D'AMMONIUM, GER, AMMONIUMCHLORID, ITAL, CLORURO DI AMMONIO, SPAN, CLORURO AMONICO

NH_4Cl , eq 53 13

White, odourless crystalline powder possessing a cooling saline taste It is permanent in the air

Solubility.—1 in 3 of Water, 1 in 55 of Alcohol (90 p c).

Medicinal Properties —Stimulating expectorant in bronchitis by inhalation, or by allowing it to dissolve slowly in the mouth in the form of lozenge or tablet, is a hepatic, gastric and intestinal stimulant, diaphoretic and diuretic In neuralgia, lumbago and migraine, in doses of 20 to 30 grains three times a day, it frequently relieves after four or five doses Useful in sciatica, gout and chronic rheumatism, in acute and chronic congestion of the liver, said to counteract the tendency to albuminoid degeneration

Recommended in advanced cases of pulmonary phthisis to facilitate expectoration —*L* '95, n 1624

Dose.—5 to 20 grains = 0.32 to 1.3 gramme

Prescribing Notes —Generally taken in solution, can be dispensed in the form of mixtures, powders, or Compressed Tablets Lemon and Chloroform make it more palatable See below, *Haustus*

Fluid Extract of Liquorice has been recommended, but many persons object to the taste of Liquorice

10 grams in a claret-glassful (3 fl oz) of cold Water, sipped frequently, allays distressing fits of coughing in bronchitis

The vapour is also largely employed in naso-pharyngeal and eustachian catarrh, various kinds of inhalers have been introduced for mixing the vapours of Hydrochloric Acid and Ammonia. In the absence of such an inhaler, heat a small quantity of the solid salt in an iron spoon or any convenient dish over a spirit lamp and inhale the fumes

Incompatibles—Alkalis and their Carbonates, alkaline earths, Lead and Silver salts

Official Preparation.—Used in the preparation of *Liquor Ammoniae Fortis*

Not Official.—Draught, Lotion and Lozenges

Foreign Pharmacopoeias—Official in Austr, Belg, Ger, Hung, Jap, Russ and Swiss (*Ammonium Chloratum*), Dan, Dutch, Norw and Swed (*Chloretum Ammonicum*), Fr (*Chlorure d'Ammonium*), Ital (*Cloruro di Ammonio*), Mex (*Cloruro de Amonio*), Port. (*Chloreto de Ammonio*), Span (*Cloruro Amonico*), US (*Ammonii Chloridum*)

Tests.—The distinguishing tests for Ammonium Chloride are the evolution of Ammonia when the salt is heated with Potassium or Sodium Hydroxide Solution and the production of a yellowish-brown coloration when a sufficiently diluted solution of the salt is treated with Potassio-mercuric Iodide (Nessler's) Solution, the depth of colour varying with the dilution of the solution, the formation of a white curdy precipitate, insoluble in Nitric Acid, but soluble in Ammonia Solution or Potassium Cyanide Solution, when Silver Nitrate Solution is added to its aqueous solution. When heated it evolves dense white fumes and volatilises completely

Neither the *BP* nor the *PG* records a volumetric method for the determination of the salt, but a process which is described below is given in the *USP*, the latter Pharmacopoeia stipulates that the salt shall contain not less than 99.5 p.c. of the pure salt

The more generally occurring impurities are Calcium, Copper, Iron, Lead, Carbonates and Sulphates. These impurities may be detected, if present, by the tests given in the small type below under the respective headings of Ammonium Oxalate Solution, Hydrogen Sulphide, Barium Nitrate or Chloride Solution, and Diluted Sulphuric Acid. The *USP* and *PG* include a separate test for Iron, which is described in the small type under the heading of Potassium Ferrocyanide Solution. It may also contain Thiocyanates. The test for the latter is carried out with Ferric Chloride Test-solution, using according to the *BP* an aqueous solution, according to the *USP* and *PG* an aqueous solution acidulated with Hydrochloric Acid.

A standard of 5 parts of Lead per 1,000,000 is suggested (*CD* '08, 1.795) and 2 parts per 1,000,000 as a standard for Arsenic

Hydrogen Sulphide—An aqueous solution (1-20 *PG*) should not be affected by Hydrogen Sulphide Solution, neither should it respond to the time-limit test for heavy metals, *USP*

Barium Nitrate or Chloride Solution.—An aqueous 1 in 20 solution should be unaffected by Barium Nitrate Solution *PG*, by Barium Chloride T.S., *USP*.

Diluted Sulphuric Acid An aqueous 1 in 20 solution should be unaffected by diluted Sulphuric Acid, *P G* and *U S P*

Ammonium Oxalate Solution—An aqueous 1 in 20 solution should be unaffected by Ammonium Oxalate Solution, *P G* and *U S P*

Potassium Ferrocyanide Solution—20 c c of a 1-20 solution should not immediately turn blue with 0.5 c c Potassium Ferrocyanide Solution, *P G*, *U S P* uses 5 drops of the reagent

Volumetric Determination—10 c c of a solution obtained by dissolving 1 gramme of Ammonium Chloride in sufficient Water to measure 100 c c, should, after the addition of 5 drops Potassium Chromate T S, require not less than 18.7 c c of Tenth-normal Volumetric Silver Nitrate Solution to produce a permanent red colour, *U S P*

Not Official

HAUSTUS AMMONII CHLORIDI.—Ammonii Chloridi, gr. xv, Tinct. Limon., m. xlv, Sp. Chloroformi, m. x, Aquæ, ad 3 iiss

LOTIO AMMONII CHLORIDI—1 oz with 1 fl oz Alcohol (90 p c.) and 10 fl oz Water. Vinegar is sometimes added, to be applied as a dressing for bruises

TROCHISCI AMMONII CHLORIDI—2 grains = 0.13 gramme, in each, with Black Currant Paste, are much used for bronchitis

Dose—2 to 4 lozenges

These have been incorporated in the *B P C*

Foreign Pharmacopœias—Official in Belg., Dutch and U S

Each lozenge contains about 2 grains of Ammonium Chloride with Black Currant Paste—*Throat*

Ammonium Chloride, 10, Extract of Glycyrrhiza, 20, Tragacanth, in fine powder, 2, Sugar, 40, in grammes, Sugar of Tolu, q s to make 100 troches—*U S P*

Trochisci Ammonii Chloridi cum Glycyrrhiza—Ammonium Chloride, 3 grains, Liquorice Extract, 3 grains in each—*Martindale*

This has been incorporated in the *B P C*

Not Official.

AMMONII IODIDUM.

AMMONIUM IODIDE

FR, IODURE D'AMMONIUM, GER, AMMONIUM JODATUM, ITAL, YODURO DI AMMONIO, SPAN, YODURO AMMONICO

A white granular deliquescent salt, or in cubical crystals, readily becoming yellow on exposure to air

The salt possesses practically no odour unless highly coloured, when a faint odour of Iodine is perceptible, and it has a sharp saline taste

When deeply coloured, it is advisable in dispensing to remove the colour by shaking it in a bottle with a piece of Ammonium Carbonate. It has been pointed out that the resulting Iodide could be decomposed by the Hydrochloric Acid of the stomach, and result in the re-formation of free Iodine, but as the quantity would generally be very small it may be disregarded

The *U S P* uses Ammonium Sulphide Test-solution for decolorising a deeply coloured salt. Sufficient of the solution to decolorise it being added to a concentrated solution, the liquid filtered and evaporated to dryness

It should be kept in well-stoppered glass bottles of a dark amber tint

Solubility—4 in 3 of Water, 1 in 3 of Alcohol (90 p c.), 3 in 4 of Glycerin.

Medicinal Properties—Similar to the Potassium Iodide, but less depressing

Dose —2 to 5 grains = 0.13 to 0.32 gramme, three times a day, but much larger doses can be given

Foreign Pharmacopœias —Official, in Fr, Mex, Port, Russ, Span Swiss and U S Not in the others

Tests —The salt answers the usual distinguishing tests for Ammonium salts, it evolves Ammonia when heated with Potassium of Sodium Hydroxide Solution, and yields a brownish yellow coloration on the addition of Potassio mercuric Iodide (Nessler's) Solution to its extremely dilute aqueous solutions. The addition of Chlorine water to its aqueous solution liberates Iodine, and a violet-coloured solution is given when this aqueous liquid is shaken with Carbon Bisulphide. Silver Nitrate Solution produces a curdy yellow precipitate, which is insoluble in Nitric Acid, practically insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution.

The more generally occurring impurities are heavy metals, *eg*, Arsenic, Copper and Lead, Barium, Iron, excess of free Iodine, and excess of Chlorides or Bromides. The presence of heavy metals is readily detected by Hydrogen Sulphide Solution, Barium by means of a solution of Potassium Sulphate. A limit of Iron is fixed by the requirement that the addition of Potassium Ferrocyanide Solution to an aqueous 1 in 150 solution of the salt should not immediately produce a blue colour. An aqueous 1 in 150 solution when shaken with 1 c.c. of Chloroform should not impart a violet colour to the chloroformic liquid, indicating the absence of excess of free Iodine. A 5 p.c. aqueous solution when acidified with Nitric Acid should neither yield a turbidity or precipitate on the addition of either Silver Nitrate or Barium Chloride Solution, indicating the absence of Chlorides and Sulphates.

In testing for excess of Chlorides or Bromides the *USP* dissolves 0.25 gramme of the salt, dried at 100° C (212° F), in 5 c.c. of Ammonia Solution, shakes with 16.9 c.c. of Deci-normal Volumetric Solution of Silver Nitrate and saturates the filtrate with 5 c.c. of Nitric Acid. The absence of a cloudiness within 10 minutes indicates the absence of more than 3 p.c. of Chlorides and Bromides.

AMMONII PHOSPHAS.

AMMONIUM PHOSPHATE

$(\text{NH}_4)_2 \text{HPO}_4$, eq 131.20

White, odourless, glistening, prismatic crystals, having a saline taste

It may be prepared by neutralisation of Orthophosphoric Acid with Ammonia Solution, the presence of the requisite amount of Ammonia being ensured by the addition of solid Ammonium Carbonate when necessary during the evaporation. It should be preserved in well-stoppered bottles.

Solubility —1 in 2 of Water, and measures 2½, insoluble in Alcohol (90 p.c.)

A salt corresponding to the official formula has been stated by some authorities to have a solubility of 1 in 0.76, the true figure, however, for the normal *BP* salt is 1 in 2, and this figure was given in the *Companion* from the 1st edition to the 15th (1890) — *C D*, '03, i 914, *P J*, '03, i 65

Medicinal Properties —Given in chronic rheumatism and in the gouty and uric acid diathesis to render the Sodium Bicarbonate more soluble, and to prevent formation of Uric Acid calculi.

Dose —5 to 20 grains = 0.32 to 1.3 grammes

Prescribing Notes. —It is given 3 or 4 times a day in Water, but should

not be prescribed in too condensed a form when tinctures form part of the mixture, on account of its sparing solubility in spirituous menstrua

Foreign Pharmacopœias—Official in Port Not in others

Tests—The distinguishing tests for Ammonium Phosphate are that it evolves Ammonia when heated with Potassium or Sodium Hydroxide Solution, its aqueous solution when sufficiently diluted yields a yellowish-brown coloration with Potassio-mercuric Iodide (Nessler's) Solution, an aqueous solution yields with Silver Ammonio-nitrate Solution a pale yellow precipitate readily soluble in Ammonia Solution and in cold diluted Nitric Acid, Ammonium Molybdate Solution when added to an aqueous solution containing an excess of Nitric Acid yields a yellow precipitate soluble in Ammonia Solution, and Magnesium Ammonio-sulphate Solution added to its aqueous ammoniacal solution affords a white crystalline precipitate soluble in dilute mineral acids. The latter test forms the basis of the *B P* method for the gravimetric determination of the salt, and it is required that when 2 grammes are precipitated with Magnesium Solution, the precipitate washed with Ammonia Solution, with an equal volume of water, suitably dried and heated to redness, shall weigh 1.680 grammes. This corresponds to 99.68 p.c. of Di-ammonium Mono-hydrogen Ortho-phosphate. It must be pointed out that unless the precautions mentioned at the commencement of the monograph on this article are observed, a salt yielding this amount of precipitate will not be obtained. Most 'commercial' samples are mixtures of Di-ammonium Mono-hydrogen Ortho-phosphate (the official salt) and Mono-ammonium Di-hydrogen Ortho-phosphate (the Acid Ammonium Phosphate).

Six commercial samples examined in the author's laboratory, when assayed by the process described in the *B P*, yielded precipitates varying in weight from 1.686 grammes to 1.818 grammes, the former being the only one of the six that approximated a salt of the Pharmacopœia formula. Greenish and F. A. Upsher Smith, from an examination of two commercial samples, neither of which yielded the official weight of precipitate, concluded (*P J* '01, 1.777) that it appeared unlikely that an absolutely normal salt could be found in commerce and accepted the second sample for the determination of its solubility, as the best that could be furnished commercially. Later (*P J* '03, 1.948), whilst confirming the investigations into the composition of commercial samples of the salt recorded in the author's paper (*C D* '02, 1.944, *P J* '03, 1.65), Greenish's further experiments have convinced him that there is no material difficulty in preparing a salt of the composition demanded by the Pharmacopœia.

The more generally occurring impurities, other than the Acid Phosphate, are Arsenic, Copper and Lead, Iron, Chlorides, and Sulphates. Of those the most important and the most likely are Arsenic and Lead, the former is readily detected by the modified Gutzeit's test, the latter by Hydrogen Sulphide Solution.

A standard of 5 parts per 1,000,000 for Arsenic and 10 parts per 1,000,000 for Lead is suggested (*C D* '08, 1.795). A 1 in 20 aqueous

solution when acidified with diluted Nitric Acid should yield no turbidity on the addition of Silver Nitrate Solution, indicating the absence of Chlorides. A solution of similar strength when acidified with Hydrochloric Acid should yield no turbidity on the addition of Barium Chloride Solution, indicating the absence of Sulphates

Not Official

AMMONIUM SALICYLATE—White, odourless, crystalline powder, or in odourless white needle shaped crystals. It may be prepared with either the natural or the physiologically pure Salicylic Acid. It should be kept in well-stoppered glass bottles of a dark amber colour and kept in a cool atmosphere. Antipyretic and antirheumatic, but is largely superseded by the Sodium salt.

Dose—5 to 15 grains = 0.32 to 1 gramme

Foreign Pharmacopœias—Official in Dutch, Russ and U.S. Not in the others.

Tests—It evolves Ammonia when warmed with Potassium or Sodium Hydroxide Solution, and its dilute aqueous solution yields with Ferric Chloride Test solution a deep violet coloration.

A small quantity of the salt warmed with concentrated Sulphuric Acid and a few c.c. of Methyl Alcohol evolves the characteristic odour of Methyl Salicylate.

Its sufficiently concentrated aqueous solution yields a white crystalline precipitate when acidified with a mineral acid, the precipitate when washed and collected should possess the melting point 156° to 157° C (312.8° to 314.6° F) of pure Salicylic Acid, and should otherwise conform to the tests for purity of the acid.

The more generally occurring impurities are heavy metals, *e.g.*, Copper and Lead, which are likely to be present in Salicylic Acid and mineral matter. The presence of heavy metals is readily detected by adding Hydrogen Sulphide Solution to the acidified, filtered, aqueous solution, impure Salicylic Acid is indicated by the melting point above described, and absence of mineral matter by the complete volatilisation of the salt on heating.

AMMONIUM VALERIANATE—Colourless, deliquescent, flat prismatic crystals, possessing a sweetish taste and a strong odour of Valerianic Acid.

It should be kept in well stoppered glass bottles of a dark amber colour and in a cool atmosphere.

Readily soluble in Water and Alcohol, soluble in Ether.

Stated to be useful in hysteria, epilepsy and neuralgia.

Dose—1 to 3 grains = 0.065 to 0.2 gramme, several times daily. Given in form of pills or in solution.

Foreign Pharmacopœias—Official in Fr., Mex., Span., Swiss and U.S. Not in the others.

Tests—The salt evolves Ammonia when heated with Potassium or Sodium Hydroxide Solution, when heated it fuses, evolving an odour of Ammonia and of Valerianic Acid, and when heated to a still higher temperature, it is completely volatilised, when warmed with diluted Sulphuric Acid the odour of Valerianic Acid is emitted. The U.S.P. requires that it shall contain not less than 98 p.c. of pure Ammonium Valerianate.

The more generally occurring impurities are heavy metals, *e.g.*, Lead and Copper and Acetates. The presence of the former is readily detected by adding Hydrogen Sulphide Solution to the acidified aqueous solution, the presence of the latter by supersaturating the aqueous solution with Ferric Chloride Test solution and filtering, the filtrate should not possess a red colour. Solutions of the salt exhibit an acid reaction to blue Litmus paper.

AMMONIUM VALERIANATUM SOLUTUM (Liquor Ammonii Pierlot), *Siems*—Valerianic Acid, 3, Extract Valerian, 2, Water, 95, Ammonium Carbonate, sufficient to neutralise.

In hysteria and epilepsy 6 to 80 drops in sweetened Water. Given in the form of the above solution in the treatment of the morphine habit.—L '01, n. 363.

AMYGDALA AMARA.**BITTER ALMOND**

FR, AMANDES AMÈRES, GER, BITTERE MANDELN, ITAL, MANDORLE AMARE; SPAN, ALMENDRA AMARGA

The ripe Seed of *Prunus Amygdalus*, Stokes, var *amara*, Baillon

Introduced only as a source of Almond Oil, and from which the commercial product is chiefly obtained

Foreign Pharmacopœias—Official in all the Foreign Pharmacopœias except Dutch, Port (Amendoas Amargas)

Not Official.—Aqua Amygdalæ Amaræ, Mistura Amygdalæ Amare, Oleum Amygdalæ Amaræ Essentiale, and Oleum Amygdalæ Persici Part of the Essential Oil, Spiritus Amygdalæ Amaræ, Syrupus Amygdalæ

Descriptive Notes.—The bitter almonds of commerce are chiefly obtained from Barbary (Morocco) and Sicily, and are distinguished by their bitter taste and the characteristic odour of the aqueous emulsion. The *USP* gives the measurement of the seed as 20 to 30 mm long, and oblong, lanceolate or ovate in form, and the *PG* as 2 cm long and 1 to 2 cm broad, and states that the seed should not have a rancid taste

Apricot and peach kernels which are imported for making an inferior almond oil are much smaller than bitter almonds, which they resemble in taste

AMYGDALÆ OLEUM. ALMOND OIL

The fixed oil expressed from the Bitter or Sweet Almond, the yield being between 40 and 45 p c

Descriptive Notes.—A clear pale yellow, odourless oily liquid possessing a bland, nutty taste. The almond oil of commerce is obtained chiefly from the bitter almond by expression, and is known in commerce as English oil. The oil formerly sold under the name of Ol Amygd Exot, but now as Ol Amygd Persic, is derived from Peach and Apricot kernels, recently Oil of Poppy Seed has been offered under the name of Peach Kernel Oil

Solubility—Only slightly soluble in Alcohol (90 p c), entirely soluble 1 in 2½ of Ether, and in all proportions of Chloroform

Medicinal Properties.—Emollient, demulcent and laxative. As an enema in impaction of fæces or obstruction of bowel, 1 to 3 pints

Dose—1 to 4 fl. drm = 3·6 to 14·2 c c

Prescribing Notes—1 fl oz of Oil, with ½ fl oz Mucilage, ¼ oz Sugar, and 6 fl oz of Distilled Water, makes a nice cough mixture

A mixture of equal parts of this Oil and Lime Water, with a small proportion of Glycerin, secreted with Lemon, has been commonly sold under the title **Glycerin** and Lime Juice

Official Preparations.—Contained in Linimentum Ammoniac, Oleum Phosphoratum, Unguentum Aquæ Rosæ, and Unguentum Catapla.

Used in preference to Olive Oil, as it makes a whiter ointment and is less liable to become rancid.

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr (Huile d'Amande), Ger (Mandelöl), Hung, Ital (Olio di Mandorle Dolci), Jap., Mex., Norw., Port., Russ., Span (Aceite de Almondas Dulces), Swed., Swiss and U S

Tests.—The distinguishing tests for Almond Oil are the specific gravity which should be from 0.915 to 0.920, the congealing point which is about -20°C (-4°F). These tests, with an Elaidin test for Peach Kernel and other fixed Oils, are as far as the official volume takes us. The above specific gravity is given by *BP* and *PG*, the *USP* gives 0.910 to 0.915 at 25°C (77°F). The *USP* and some of the recent editions of the Continental Pharmacopœias include figures for the Saponification value and Iodine absorption. The Saponification value is the number of milligrammes of Potassium Hydroxide Solution required to neutralise 1 gramme of the oil. The Iodine absorption is the percentage of Iodine which the oil absorbs. The Austr. Ph. requires a Saponification value of 190–195 and an Iodine absorption of 94–100 p.c., the Belgian Ph., a Saponification value of not less than 185, and an Iodine absorption of 92–102 p.c. The *USP*, a Saponification value of 191–200 and an Iodine absorption of not less than 95 nor more than 100, the Dutch Ph. omits the Saponification value, but gives an Iodine absorption of not less than 95 nor more than 101. Twelve good commercial samples examined in the author's laboratory showed a Saponification value of 190.4 to 196.4, and an Iodine absorption of 95.3 to 100.3.

The more generally occurring impurities are Peach and Apricot Kernel Oils (which much resemble Almond Oil, and are known commercially as Ol. Amygdalæ Persicæ), and other fixed oils, e.g., Arachis, Cotton Seed, Olive and Sesame. The Elaidin test for these, which is given in small type below under the heading of Nitric Acid, is common to *BP*, *USP* and *PG*. The two latter Pharmacopœias, moreover, include an additional, almost identical, test for the detection of fixed oils other than Almond Oil, which is given in small type below under the heading of Saponification, and which depends upon the production of a clear solution when the saponified alcoholic solution is diluted with Distilled Water, the Oleic Acid obtained from this solution on acidification, when washed and clarified, is required to remain liquid at 15.5°C (60°F), and when mixed with an equal volume of Alcohol of the respective pharmacopœial strengths to yield a clear solution which does not deposit at 15°C (60°F) nor become turbid on a further addition of one volume of Alcohol.

Nitric Acid.—If 2 c.c. of the Oil be well shaken with 1 c.c. of fuming Nitric Acid and 1 c.c. of water, a whitish, not brownish, red mixture should be formed which, after standing for 6 hours at about 50°F (10°C) (for 2 or at most 6 hours *PG*, for some hours *USP*), should separate into a solid white mass and a nearly colourless liquid, *BP*, *PG*, and *USP*. The *USP* states that the fixed Oils of Peach and Apricot kernels give a red colour and Sesame and Cotton Seed Oils give a brown colour.

Saponification.—If 10 c.c. of the Oil be mixed with 15 c.c. of a Sodium Hydroxide Solution (1-G, *USP*, sp. gr. 1.168 to 1.172, *PG*), and 10 c.c. of Alcohol, and the mixture be allowed to stand (at a temperature of 35° to 40°C (95° to 104°F)) with occasional agitation, (*USP*) until it becomes clear, and if then diluted with 100 c.c. of Water, the clear solution thus obtained upon the

subsequent addition of an excess of Hydrochloric Acid will set free a layer of Oleic Acid. This when separated from the aqueous liquid, washed with warm Water, and clarified by heating on a water-bath will remain clear if cooled to 15° C (59° F), *P G* and *U S P*. 1 volume of this Oleic Acid when mixed with 1 volume of Alcohol should yield a clear solution which at 15° C (59° F) should not deposit any fatty acids, nor become turbid upon the further addition of 1 volume of Alcohol, *P G* and *U S P*.

Iodine Absorption—If 0.3 gramme of expressed Oil of Almonds be dissolved in 10 c.c. of Chloroform in a 250 c.c. flask and 25 c.c. of a mixture of equal volumes of Alcoholic Iodine T.S. and Alcoholic Mercuric Chloride T.S. added, and if, after standing for four hours protected from light, 20 c.c. of Potassium Iodide T.S. be introduced and the mixture diluted with 50 c.c. of Water, on titrating the excess of Iodine with V.S. of Tenth-normal Sodium Thiosulphate an Iodine value of not less than 95 nor more than 100 should be obtained, *U S P*. The *P G* test uses 0.5 gramme of Oil, and directs the addition of 1.5 grammes of Potassium Iodide and 100 c.c. of Water before titration.

Not Official.

AQUA AMYGDALÆ AMARÆ—Prepared by crushing Bitter Almonds and expressing the fixed Oil, and then distilling the residual cake with Water so that it shall contain the proper quantity of Hydrocyanic Acid ordered in any particular Pharmacopœia.

Ph. Ger. maximum single dose, 2 grammes, maximum daily dose, 6 grammes.

Foreign Pharmacopœias—Official in the following, the percentage of Hydrocyanic Acid also given: Dan (Conc), 0.1 p.c., (Dil) 0.005 p.c., Ger, Hung, Ital, Jap, Russ and Swed, 0.1 p.c., Norw, 0.1 p.c., Port, not standardised, U.S., not standardised, 1 Volatile Oil in 1000, Austr, Belg and Swiss use Laurocerasi Water when Bitter Almond Water is prescribed. Not in the others.

The *Brussels Conference* adopts a strength of 0.1 p.c. for Aqua Amygdalæ Amaræ.

MISTURA AMYGDALÆ AMARÆ.—Made in the same proportions as Mistura Amygdalæ.

Useful in cough, and as a lotion to allay itching of the skin. It was a favourite vehicle for giving Tartarated Antimony, in doses of $\frac{1}{2}$ grain = 0.008 gramme, as a sedative expectorant in the first stage of acute bronchitis or pneumonia. The mixture contains a variable amount of Hydrocyanic Acid.

Dose— $\frac{1}{2}$ to 1½ fl. oz = 14.2 to 42.6 c.c.

Mistura Amygdalæ Amaræ—Bitter Almonds, 8, Distilled Water, q.s. to produce 100.—*B P C*.

OLEUM AMYGDALÆ AMARÆ ESSENTIALE—A clear, colourless or pale yellowish, highly refractive liquid, with a characteristic odour, and a bitter and somewhat burning taste. Obtained from Bitter Almonds by macerating with Water the cake from which the fixed Oil has been expressed, and subsequent distillation.

It should be kept in well-stoppered glass bottles of a dark amber colour, and as far as possible from contact with air.

Chiefly used as a flavouring agent, when the oil 'sine Acido Hydrocyanico' should be employed.

Oil Amygdal Essent Persic is prepared by a similar process to Bitter Almond Oil, from the kernels of the Apricot and Peach.

Solubility—Sparsely in Water; mixes in all proportions with Alcohol (90 p.c.) and Ether.

Foreign Pharmacopœias—Official in Belg (Aldehydum Benzœicum), Fr. Mex (Acete Volatil de Almendras Amargras), Port, Span and U.S. U.S. has also Benzaldehyde. Not in the others.

U.S. has also Spiritus Amygdalæ Amaræ, 1 in 100.

Tests.—The distinguishing tests for Bitter Almond Oil are the strong and distinctive odour, the high refraction, the specific gravity which after removal of Hydrocyanic Acid should be from 1.045 to 1.050, the boiling point, which should be 179° to 180° C (354.2° to 356° F), and its optical inactivity. The Oil is converted into a crystalline compound when shaken with a saturated Solution of Sodium Bisulphite, and this reaction may be utilised as a means for its quantitative determination, the non aldehydic constituents can be measured or weighed. The *U S P* process is essentially as follows.—A measured quantity of 10 c c of purified Kerosene is introduced into a tared 150 c c flask and the exact weight recorded, 12 drops of the Oil are then added and the weight again recorded, 20 c c of Distilled Water and 6 drops of Rosolic Acid Test solution are added and the mixture exactly neutralised with Tenth-normal Volumetric Sodium Hydroxide Solution, agitating the flask thoroughly. A 1 in 5 aqueous solution of Sodium Sulphite alternated with Half-normal Volumetric Hydrochloric Acid Solution is added until 10 c c of the Sodium Sulphite Solution have been added, and sufficient Half-normal Volumetric Hydrochloric Acid Solution to maintain the neutrality of the mixture, after the addition of a few drops of Rosolic Acid Test-solution, the flask is agitated frequently, allowed to stand for 2 hours to ensure a permanent condition of neutrality, and the number of c c of Half normal Volumetric Hydrochloric Acid Solution required noted. A blank test is carried out alongside of the determination, the number of c c of Half-normal Volumetric Hydrochloric Acid Solution is noted, the number of c c used in the latter is subtracted from that required in the former, the difference is multiplied first by 0.0526 and the product by 100 and divided by the weight of Oil taken yields a percentage of Benzaldehyde present in the sample.

The more generally occurring impurities are acidity, Hydrocyanic Acid, artificial Benzaldehyde, and Nitrobenzine. Acidity may arise from atmospheric oxidation of Benzaldehyde to Benzoic Acid, and may be determined with Deci-normal Volumetric Sodium Hydroxide Solution. If the Oil contains a crystalline deposit of acid it should not be used.

Hydrocyanic Acid can be detected by shaking 10 or 15 drops of the Oil with 2 to 3 drops of Potassium or Sodium Hydroxide Solution, adding a few drops of Ferrous Sulphate Solution containing a drop or two of Test-solution of Ferric Chloride, warming and slightly acidifying with dilute Hydrochloric Acid, when a blue precipitate will be produced if this acid be present.

Hydrocyanic Acid may be estimated volumetrically by weighing 1 gramme into a small flask, adding a sufficiency of freshly precipitated (Chloride free) Magnesium Hydroxide and Water, several drops of Potassium Chromate Solution and Deci-normal Volumetric Solution of Silver Nitrate until a permanent red coloration is produced. The oil generally employed in this country is free from this acid, but *U S P* allows not less than 2 p c nor more than 4 p c. The presence of Chlorine compounds is generally held to be indicative of artificial Benzaldehyde, but the failure to find them does not necessarily imply that the specimen is free from 'artificial Benzaldehyde' as the latter is now produced commercially of very fine quality.

Chlorine compounds are examined for as follows.—A folded strip of filter paper saturated with oil is placed in a small porcelain capsule standing within a larger one and ignited. A large beaker, the sides of which have been moistened with Water, or preferably containing filter paper (free from Chlorides) moistened with Water, is inverted over the burning Oil. The products of combustion condense on the moistened surface of the beaker or on the filter paper, and can be washed on to a filter with a little Distilled Water. The filtrate should show no turbidity on the addition of Silver Nitrate Solution, or if a turbidity is produced it should disappear on warming. The presence of Nitrobenzene (now an unlikely impurity) may be detected by diluting the Oil with 20 times its volume of Alcohol (90 p c), diluting the solution with Water until a turbidity is produced, adding Zinc and Sulphuric Acid, allowing the solution to remain at rest several hours, filtering, evaporating the Alcohol, and treating the remaining solution with a drop of Potassium Bichromate Solution. The production of a violet colour indicates the presence of Aniline.

SPIRITUS AMYGDALÆ AMARÆ—Oil of Bitter Almond, 1, Alcohol (95 p c), 80, Distilled Water, q.s. to make 100.—*U. S. P.*

This spirit contains a trace of Hydrocyanic Acid and should always be used with care

This has been incorporated in the *B P C* as follows —

Oil of Bitter Almond (sine Acid Hydrocyanic), 1, Alcohol (90 p c), 85,
Distilled Water, *q s* to produce 100

SYRUPUS AMYGDALÆ—Spirit of Bitter Almond, 1, Orange Flower
Water, 10, Syrup, *q s* to make 100—*U S P*

This has been incorporated in the *B P C*

AMYGDALA DULCIS.

SWEET ALMOND

FR, AMANDES DOUCES, GER, SÜSSE MANDFELN, ITAL, MANDORLE DOLCI,
SPAN, ALMENDRO DULCE

The ripe Seed of *Prunus Amygdalus*, Stokes, var *dulcis*, Baillon
Commonly known as the Jordan Almond

Medicinal Properties.—Demulcent and nutrient Biscuits are made of Jordan and Valencia Almonds for diabetic patients, as a substitute for bread or starchy food Almonds do not contain starch

The Mistura Amygdalæ is a good vehicle for cough medicines, and for suspending insoluble powders

Official Preparations—Mistura Amygdalæ and Pulvis Amygdalæ Compositus

Foreign Pharmacopœias—Official in all, Fr, Ital, Mex (Almendra Dulce), Port (Amendoas Doces), Span

Descriptive Notes.—Almonds are met with in commerce either in the shell or endocarp, or in the form of seed only Sweet Almonds vary in size and shape, those of Valencia and Sicily being broad and flattened, those of Barbary irregular in shape owing to two seeds often occurring in one endocarp and becoming misshapen by pressure The Valencia Almonds are usually free from Bitter Almonds, those of Barbary and Sicily are often mixed with them The Jordan Almond, which is alone official, is imported from Malaga in Spain It is longer, narrower and more convex in proportion than the other varieties, being 2·5 to 3 cm (about an inch) in length, and about 1·25 cm ($\frac{1}{2}$ inch) in width and 0·5 cm ($\frac{1}{4}$ inch) in thickness It has a bland taste and triturated with Water forms a white emulsion without any marked odour, indicating the absence of Bitter Almonds For forming emulsions, Almonds are first deprived of their skins or 'blanched' by dipping them in boiling Water for a minute or two, when slight pressure between the fingers will separate the testa The variety official in the *U S P* appears to be the Valencia Almond as it is stated to be broader than the Bitter Almond The variety official in Germany is stated to be unsymmetrically ovate and flattened, 2·25 cm long and 1·5 cm broad, and at the rounded end more than 1 cm broad. This appears to indicate the Sicilian or Valencia Almonds, since the Jordan Almonds are rarely more than 1·25 cm broad.

Preparations

MISTURA AMYGDALÆ. ALMOND MIXTURE

Compound Powder of Almonds, 1, Distilled Water, 8

Rub the powder to a smooth cream with a suitable quantity of Water, and add gradually the remainder, strain

Dose — $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

Foreign Pharmacopœias — Official as Emulsio in Austr, Dan, Fr, Hung, Ital, Norw, Port and Swed US Emulsum A, there is much variation in the proportions Swed has also Emulsio Hydrocyanata, Ayrngdalin, 1, Almond Emulsion, 80 Not in the others Belg, Fr, Ger and Swiss have Syrups

PULVIS AMYGDALÆ COMPOSITUS — COMPOUND POWDER OF ALMONDS

Sweet Almonds, 8, Powdered Refined Sugar, 4, Powdered Gum Acacia, 1 Remove the skins of the Almonds after softening them in Water, and dry the Almonds by a cloth and exposure to the air until brittle (*Comp* 1894), so that they will rub to a paste which is not too moist and with which the Sugar and Gum, previously mixed, can be incorporated to form a moderately coarse powder

Dose — 60 to 120 grains = 4 to 8 grammes

AMYL NITRIS.

AMYL NITRITE

FR., AZOTITE D'AMYLE, GER, AMYLNITRIT, ITAL, NITRITO D'AMILE, SPAN, NITRITO DE AMILO

A pale yellow, volatile liquid, with a characteristic ethereal odour and pungent aromatic taste It consists principally of Iso-Amyl Nitrite, $C_5H_{11}NO_2$, eq 116 25, which is present in variable quantity, together with other Nitrites It may be produced by the action of Nitrous Acid upon that fraction of the higher Alcohols distilling between 127 7° and 132 2° C (262° and 272° F) Should be stored in well-stoppered glass bottles of an amber colour, or preferably in glass capsules of a dark amber tint

Solubility — Insoluble in Water Soluble in Alcohol (90 p c), Ether and Chloroform

Medicinal Properties. — Antispasmodic Very useful in angina pectoris, aneurismal pain, hæmoptysis, dyspnoea of bronchitis and spasmodic asthma, has been used with advantage in epilepsy, in trifacial neuralgia, in migraine and sea-sickness, and hemicrania, if these conditions be accompanied by facial pallor, also in laryngeal spasm, in hepatic, intestinal and renal colic, in spasmodic forms of dysmenorrhœa and in eclampsia, a restorative in cardiac failure from Chloroform or Nitrous Oxide anæsthesia or other cause, has been found useful as an antidote to Strychnine

In angina, where a rapid fall of arterial tension is required, the

B.P. Amyl Nitrite contains Iso-butyl Nitrite, is best, but in other cases, such as in the disease, when the prolonged action is required, pure Nitrite is more effective. As some persons are peculiarly susceptible to its action, its use demands caution.

A description of 77 cases of pneumonia treated by the inhalation of large doses—*B.M.J.* '95, ii 96, *T.G.* '96, 49.

Promptly effective and safer than Morphine in hæmoptysis, and can be used at earliest possible moment by patient himself—*L.* '08, i 565.

The most efficient and expeditious remedy in hæmoptysis—*L.* '07, i 679, *T.G.* '07, 323, *L.* '06, ii 1685, '07, i 939, '08, i 427, 504.

Successful in postpartum hæmorrhage (*B.M.J.* '06, ii 1125), and in menorrhagia—*L.* '08, ii 418.

Inhalations have been recommended (*L.* '05, i 800) in deep-seated hæmorrhage. It has also been used with success in tuberculous hæmoptysis.

Dose—For inhalation, the vapour of 2 to 5 minims = 0.12 to 0.3 cc.

Prescribing Notes—It can be obtained in small glass capsules covered with cotton wool and silk, each containing from 2 to 5 minims. The covered capsule is placed in a handkerchief, then carefully broken across, and the escaping vapour is inhaled.

In mixtures to be swallowed, dose, $\frac{1}{2}$ to 1 minim dissolved in Alcohol (90 p.c.) and diffused through Water by means of Tragacanth (in powder) 2 grains to the fl. oz., to be used with caution.

Should be handled carefully, as even smelling it causes violent flushings.

Not Official—Iso-butyl Nitrite, Tertiary Amyl Nitrite, Amyl Valerianate and Mistura Amyl Nitritus.

Foreign Pharmacopœias—Official in Austr., Dutch, Ger., Jap., Russ. and Swed., sp. gr. 0.870 to 0.880, boils at 97° to 99° C., Belg., sp. gr. 0.870 to 0.900, boils at 99° C., Fr., sp. gr. 0.88, boils at 96° to 99° C., Mex. (Ester Amilnitroso), sp. gr. 0.877, boils at 95° C., Ger., Russ. and Swed., boils at 97° to 99° C., Hung., sp. gr. 0.900, boils at 96° to 99° C., Ital. (Nitrito amilico), sp. gr. 0.87 to 0.89, boils at 97° to 99° C., Swiss, sp. gr. 0.870 to 0.900, boils at 97° to 99° C., U.S., sp. gr. 0.865 to 0.875 at 25° C. (77° F.).

Tests.—The distinguishing tests for Amyl Nitrite are its odour; the peculiar flushing of the face and strange sensation of fullness in the head produced by its inhalation, the specific gravity which should be from 0.870 to 0.880, the temperature [90° to 100° C. (194° to 212° F.)] at which the greater portion of the liquid passes over when distilled with the bulb of the thermometer not touching the surface of the residual liquid, the production of Iso-valerianate when it is dropped gradually on to fused Potassium Hydroxide. The *B.P.* and the *P.G.* give the specific gravity as 0.870 to 0.880; the *U.S.P.* gives 0.865 to 0.875 at 25° C. (77° F.). The *B.P.* requires that 70 p.c. should pass over between the temperatures indicated above, the *P.G.* states that it boils at 97° to 99° C. (206.6° to 210.2° F.), the *U.S.P.* has deleted the boiling point. The *B.P.* requires that only a pale yellow colour shall be produced in the aqueous liquid when the Nitrite is shaken with an equal volume of Potassium Hydroxide Solution, the *U.S.P.* and *P.G.* require that Silver Nitrate Solution should not turn brown or black. The *U.S.P.* and the *P.G.* both adopt a test for the limit of acidity, which is given below under that heading. The *B.P.* requires it to yield not less than 6 times its volume of Nitric Oxide gas, the *U.S.P.* requires it to

contain about 80 p c of Amyl Nitrite, chiefly Iso-amyl Nitrite, as determined gasometrically by the process given below under the heading of Gasometric Determination. When a measured quantity of 5 cc of a 5 p c solution of the Nitrite in Alcohol (90 p c) is intermittently shaken for 5 minutes in a nitrometer containing saturated brine solution, with 5 cc of strong Potassium Iodide Solution and 5 cc of Diluted Sulphuric Acid, and the level of the liquid in the two limbs of the nitrometer adjusted to the same level, it should yield a volume of gas not less than 30 cc, adjusted to the normal temperature and pressure. The number of cc of gas evolved multiplied by 5 (4.98) gives the weight in milligrammes of Amyl Nitrite in the quantity operated upon. The operation may be conducted in two parts, the Potassium Iodide Solution being first run into the nitrometer, followed by the Diluted Sulphuric Acid. The measure of the gas first liberated affords a criterion of the acidity of the sample. The *USP* neutralises any acidity by first treating the Amyl Nitrite with Potassium Bicarbonate and weighs the Nitrite instead of measuring it. The temperature at which the readings are taken is 25° C (77° F) and a correction is made for each degree above or below. A correction is also made for the barometric pressure above or below 760 mm of Mercury.

The more generally occurring impurities are Water and Aldehyde. All three Pharmacopœias adopt the method of cooling the specimen to the temperature of melting ice, in testing for Water, if water be absent it will remain clear. Aldehyde in the *BP* is examined for by Potassium Hydroxide Solution, no quantities being given, in the *USP* and *PG* by Silver Nitrate Solution rendered slightly ammoniacal by the addition of a few drops of Ammonia Solution, both Pharmacopœias carefully stating the quantities to be employed in the test, the *USP* uses Alcohol (94.9 p c) for diluting, the *PG* Absolute Alcohol.

Acidity—5 cc Amyl Nitrite should not overcome the alkaline reaction of 0.1 cc Solution of Ammonia with 1 cc of Water, *PG*. If 5 cc be agitated with a mixture of 1 cc Normal Potassium Hydroxide Solution and 10 cc of Water with a drop of Phenolphthalein T S, the red tint of the aqueous layer should still be perceptible, *USP*.

Silver Nitrate Solution—1 cc Amyl Nitrite should not turn brown or black a mixture of 1.5 cc Silver Nitrate Solution, 1.5 cc Alcohol and a few drops of Solution of Ammonia on gently warming (test for Aldehyde), *PG* and *USP*. The *PG* uses Absolute Alcohol in the place of Alcohol.

Gasometric Determination—Transfer about 3 cc of Amyl Nitrite, which has been previously shaken with 0.5 gramme of Potassium Bicarbonate and carefully decanted, to a tared 100 cc measuring flask, and weigh it accurately. Add sufficient Alcohol (94.9 p c) to bring the volume to exactly 100 cc and mix thoroughly. Introduce into a nitrometer exactly 10 cc of the alcoholic solution, followed by 10 cc of Potassium Iodide T S, and afterwards by 10 cc of Volumetric Sulphuric Acid Solution. When the volume of gas has become constant (within 30 to 60 minutes), note the volume of gas collected. Multiply this volume in cc by 4.8, and divide the product by the original weight of the Amyl Nitrite, the quotient will represent the percentage of Amyl Nitrite in the liquid at standard temperature and pressure. Correction for temperature and pressure. The temperature correction is one-third of one per cent of the total percentage just found for each degree, additive if the temperature is below, and subtractive if it is above 25° C (77° F). The barometric correction is four-

thirtieths of 1 p c for each mm, additive if it is above, and subtractive if it is below 760, *U S P*

For comments see above in large type

Not Official.

MISTURA AMYL NITRITIS—Amyl Nitrite, 2; Alcohol (90 p c), 16; mix and add to Powdered Tragacanth, 1, contained in a dry phial, then add gradually Distilled Water to 240, shake well **Dose**—1 or 2 dr (3 5 to 7 c c)
—*Martindale*

Amyl Nitrite, 1½ minims, Alcohol, 12 minims, Tragacanth, 1 grain; Syrup, 80 minims, Distilled Water, to 4 fl dr—*B P C*

ISO-BUTYL NITRITE—Its action and uses are similar to those of Amyl Nitrite

TERTIARY AMYL NITRITE (Bertoni's Ether)—Prepared from tertiary Amylic Alcohol (Amylene Hydrate) It possesses all the properties of the official Nitrite, but it can be taken in larger quantities without danger, and it does not cause flushing of the face

The tertiary Nitrites have a more powerful influence generally than the secondary or primary

Dose—5 drops on sugar, or in capsules

AMYL VALERIANAS—A colourless liquid, possessing a strong fruity odour Sedative and antispasmodic

Dose.—2 to 3 minims = 0 13 to 0 2 c c. in capsules

Not Official.

AMYLENE HYDRATE.

TERTIARY AMYLIC ALCOHOL DIMETHYL-ETHYL CARBINOL

$C_5H_{12}O$, eq 87 43

A clear, colourless, oily liquid, with a strong characteristic odour and taste

Solubility—1 in 8 (or rather less) of Water, in all proportions of Alcohol (90 p c).

Medicinal Properties.—Hypnotic Has no unpleasant after-effects, and its taste is less objectionable than that of Paraldehyde Successful in mania (especially morphinomaniac, *M A* '94, 426), delirium tremens, and in severe forms of epilepsy where bromides are found useless

Recommended where hypnotics are required for a long period.—*Y B T*, '94, 74.

1½ grammes Amylene Hydrate repeated in two hours gave six hours' uninterrupted sleep in a case recovering from the morphia habit—*L*, '01, 11 366.

Dose—30 to 60 minims = 1 8 to 3 6 c c

Prescribing Notes—Dissolved in Water or Alcohol (90 p c), also given in capsules, sometimes given as an enema

Cannot be employed subcutaneously owing to pain produced.—*B. M. J. E.*, '94, 11 64

Foreign Pharmacopœias—Official in Dan, Ger, Norw. and Swiss (Amylenum Hydratum) Not in the others

Tests—It should possess a specific gravity of 0 812 to 0 814 at 15° C (point of 99° C (210 2° F) to 103° C (217 4° F) 1 c c dissolved in 20 c c. of water should not within 10 minutes either decolorise 2 drops of a 1 in 1000 Potassium Permanganate Solution, indicating the absence of Ethyl or Amyl Alcohol, or blacken Silver Nitrate Solution at 100° C (212° F.), indicating the absence of Aldehyde.

AMYLUM.

STARCH

FR, AMIDON DE BLE, GER, WEIZENSTARKE, ITAL, AMIDO, SPAN, ALMIDON

A white, odourless, tasteless, impalpable powder, or irregular, angular or columnar masses, procured from the Fruits or grains of wheat, *Triticum sativum*, Lam, maize, *Zea Mays*, L, and rice, *Oryza sativa*, L

Medicinal Properties—Protective, absorbent A good application to the skin when irritable or inflamed, or in trivial burns It has been given in powder for diarrhoea, and is a very good antidote for Iodine poisoning, followed by an emetic Mucilage of Starch, 1 in 40, is useful as a basis for enemas In the form of Violet Powder it is useful to prevent the chafing and excoriation of the skin of infants Glycerin of Starch is a good application for chilblains and chapped hands, and as a protective in certain skin diseases

Official Preparations.—Glycerinum Amyli Used in the preparation of Pulvis Tragacanthæ Compositus

Not Official—Mucilage of Starch, Test Solution of Starch and Pulvis Viola

Foreign Pharmacopœias—Official in Austr, Wheat and Rice, Belg, Arrowroot, Maize, Potato, Rice and Wheat, Fr, Ger, Hung, Ital, Mex, Norw, Port, Russ, Span and Swed, all Wheat Starch Dan, Arrowroot and Wheat, Dutch, Arrowroot, Potato, Rice and Wheat, Jap, Katakuri, Kuzu and Potato, Port allows several other Starches, Swiss, Rice and Wheat Starch, U S, Maize Starch Fr has also Potato Starch

Descriptive Notes—Starch is met with in commerce in irregular columnar or pseudocrystalline masses, which may be white or coloured slightly blue, or in powder The starches most commonly used for food are those of Maize, Rice and Potato, although a number of others are met with, descriptions of which with measurements and excellent illustrations are given in Greenish's *Anatomical Atlas*, pl 1-ix, pp 6-22, and in Tschurch and Oesterle's *Anatomische Atlas* (1900)

The official Starch includes those of Wheat, Maize and Rice, and may be either in the columnar form or in powder, but should be white and inodorous Wheat Starch is described as consisting of large and small granules, the larger being lens-shaped and faintly striated concentrically, with a nearly central hilum, Maize Starch as frequently polygonal, more uniform in size, and somewhat smaller than the large granules of Wheat Starch, with a distinct hilum but no striæ, and Rice Starch as consisting of extremely minute granules nearly uniform in size, polygonal, and without evident hilum or striæ Potato Starch, by reason of its cheapness, is employed for many technical purposes and may be expected to occur as an adulteration of other Starches It should be noted that the lenticular grains of Wheat Starch when seen edgewise under the microscope appear elliptical or almost linear, and might be mistaken for the flattened grains of Zingiberaceous Starches, that a hilum is rarely evident

Arrowroot, obtained from *Maranta arundinacea*, was formerly

official, like all other Starches it is apt to absorb the odour and flavour of any drugs or perfumes near which it is placed and should consequently be kept in jars or tins

The commercial Starches may to some extent be distinguished by the different tints assumed by them when placed under a bell glass around a crystal of Iodine, and differ also in the character of the jelly they form with the same proportion of Water or Glycerin. In the *P G* only the Starch of Wheat is official, the measurements of the grains are given as 0.015 to 0.045 mm broad, and 0.002 to 0.008 mm in diameter. In the *USP* Maize Starch is alone official and the diameter of the grain is given as 0.010 to 0.025 mm.

Tests.—The distinguishing tests for Starch are the production of a translucent colourless gelatinous solution when boiled with water; the production of a deep blue colour when Iodine Solution is added to this solution when cold, the ready hydrolysis with the formation of products having a strong reducing action upon Fehling's Solution (Potassio-cupric Tartrate Solution) when the gelatinised solution is treated at a temperature of 100° F (37.7° C), with an active solution of an amylolytic enzyme.

The more generally occurring impurities are free alkali and an excessive amount of mineral matter. The *BP* includes as the official Starches, Wheat, Rice and Maize, the *USP*, Maize Starch, the *P G*, Wheat Starch only. Both *BP* and *USP* requires that when triturated with cold Water it shall yield a mixture having neither an acid nor an alkaline reaction to test papers, the *P G* stipulates that 1 part by weight of Starch boiled with 50 parts by weight of Water shall yield a mucilage which does not alter Litmus paper. Neutral Starch is, however, seldom obtained, it is, as a rule, faintly alkaline.

The *BP* makes no reference to the amount of ash permissible, the *USP* and *P G* state that not more than 1 p.c. of residue shall remain after complete incineration.

The *USP* requires that when freed from Water by careful drying in a current of warm air, Starch should show not less than 95 p.c. of hydrolysable carbohydrates.

Preparation.

GLYCERINUM AMYLI. GLYCERIN OF STARCH

Starch, 1, Glycerin, 6½, Distilled Water, 1½; stir them together whilst sufficient heat is applied to burst the Starch granules, and form a homogeneous mass.

The operation should be conducted as quickly as possible, to avoid excessive loss of Water, and to prevent carbonisation from overheating, the use of an oil-bath is recommended.

This formula has been altered in each successive edition of *B.P.* In 1867 the formula was Starch 1, Glycerin 8, in 1885, Starch 1, Glycerin 5, Distilled Water 3, and the proportions are now as that given above.

Foreign Pharmacopœias—Official in Belg, Starch 10, Water 15, Glycerin 90, Fr. (Glycéré d'Amidon), Starch 1, Water 1, Glycerin 18; Ital (Glicerolato d'Amido), Starch 7, Water 8, Glycerin 90; Mex. (Glicera de Almidon), Starch 2.4, Glycerin 80; Port. (Glicerado Com-

mum), Starch 1, Water 2, Glycerin 17, Span (Glicerolado de Almidon), Starch 1, Water 2, Glycerin 8, US (Glyceritum Amyli), Starch 1, Water 1, Glycerin 8 The following are called Unguentum Glycerini, Austr, Starch 1, Water 2, Glycerin 10, Dan, Starch 3, Water 3, Glycerin 14, Ger and Jap, Starch 10, Water 15, Glycerin 90, Hung, Tragacanth 1, Alcohol 5, Glycerin 50 (no Starch), Norw and Swed, Starch 1, Glycerin 15½, Russ, Starch 1, Water 1, Glycerin 14, Swiss, Starch 7, Water 7, Glycerin 93 All by weight

Not Official

MUCILAGE OF STARCH—A recently prepared solution containing approximately 2 p c w/v of Starch prepared by first rubbing 1 gramme of Starch to a smooth paste with Water, and adding a further sufficient quantity of Water to produce 50 c c, after boiling for a few minutes the mixture is cooled

PULVIS VIOLÆ—Orris Rhizome, in fine powder, 12 lb, Oil of Bergamot, 1 fl oz, Otto of Rose, 144 minims, Tincture of Musk (Squire), 8½ fl oz, Starch, in powder, 112 lb—*Squire*

Orris Rhizome, in fine powder, 12 50, Oil of Bergamot, 0 25, Oil of Neroli, 0 02, Starch, in powder, q s to produce 100—*B P C*

TEST SOLUTION OF STARCH—*See Appendix*

Not Official

AMYLUM IODATUM

Iodine, 5, Starch, 95, Distilled Water, q s Triturate the Iodine with a little Distilled Water, add the Starch gradually, and continue triturating until the compound assumes a uniform blue colour approaching black Dry at a temperature not exceeding 40° C (104° F) and rub it to a fine powder

This has been incorporated in the *B P C*

A teaspoonful thrice daily for lupus erythematosus—*B M J '80*, 1 652

Not Official

ANALGEN.

BENZANALGEN, QUINALGEN, ORTHOETHOXY-ANA-MONOBENZOYLAMIDOCINOLIN.

A white crystalline powder, inodorous and tasteless

This is similar in chemical composition and properties to Phenacetin, but with the Phenol ring replaced by the Quinoline ring

Solubility—Insoluble in Water, sparingly soluble in cold, more so in hot Alcohol, fairly soluble in Chloroform, almost insoluble in Ether

Medicinal Properties—Has been recommended in neuralgia, hemicrania, and bronchitic asthma, but it is not without unpleasant effects, the urine is frequently coloured red, toxic action and dangers—*B M J '98*, 11 1055

It has given relief in sciatica—*M A '94*, 9, *B M J E '93*, 11 87, *M P '94*, 621, *L '97*, 1 1227

Dose—7½ to 15 grains = 0.5 to 1 gramme

Prescribing Notes.—*Usually given in cachets, or Compressed Tablets.*

Tests—Analgen melts at a temperature of 208° C (406.4° F), and should leave no residue on ignition The cold saturated aqueous solution should yield a yellow coloration with Ferric Chloride Test solution It dissolves in cold concentrated Sulphuric Acid, forming a bright yellow coloured liquid, and on dilution with Water a lemon yellow coloured precipitate is thrown down The cold saturated aqueous solution reduces Silver Nitrate Solution in the cold or on warming.

ANETHI FRUCTUS.

DILL FRUIT.

FR, ANETH, GER, DILLSAMEN, ITAL, ANETO, SPAN., ENELDO.

The dried ripe Fruit of the *Peucedanum graveolens*, Benth and Hook f.

Cultivated in Britain or imported from Central and Southern Europe

Descriptive Notes.—Two varieties of Dill are met with in commerce, viz, the European and the Indian. The former only is official. The fruits are oval, flat, about $\frac{1}{8}$ inch (4 mm) long, $\frac{1}{8}$ inch (2 to 3 mm) broad, brown, with the two outer ridges developed into a pale marginal wing. A transverse section shows 6 vittæ. The odour and taste are aromatic.

The Indian Dill is narrower, more elliptical, more convex, and of a greyish tint, and is usually not so free from fruit-stalks as the European drug. The oil obtained from it differs in containing Dill Apioi, and less Carvone, as well as in chemical and physical characters, from that of European Dill. The plant yielding it is *Peucedanum Soua*, Benth and Hook f, considered by some botanists as only a variety of *P. graveolens*, but the *BP* description excludes the use of the Indian drug. The Dill Apioi separates during distillation, forming a layer at the bottom of the receiver.

Medicinal Properties.—Stomachic and carminative; chiefly given to children in cases of flatulency, sometimes given with Sodium Bicarbonate, the taste of which it covers well.

Official Preparations.—Aqua Anethi and Oleum Anethi.

Foreign Pharmacopœias.—Official in Mex (Eneldo), Port (Endro). Not in the others.

AQUA ANETHI. DILL WATER

Dill Fruit, 1, Water, 20, distil, 10. (1 in 10)

Not in the other Pharmacopœias

Dose.— $\frac{1}{2}$ to 1 fl oz. = 14.2 to 28.4 c.c., for children, 60 minims = 3.6 c.c.

OLEUM ANETHI. OIL OF DILL

A pale yellow, thin, oily liquid having a characteristic odour resembling Caraway, and possessing at first a sweetish and aromatic and subsequently a sharp, burning taste.

The Oil distilled from Dill Fruit.

Yield 3 to 4 p.c.

It darkens in colour on exposure to air and light and should be preserved in well-stoppered bottles of a dark amber tint. It contains about 40 to 60 p.c. of Carvone, and a terpene Limonene, but no Anethol.

Solubility.—Readily soluble in Alcohol and Ether.

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Not in the other Pharmacopœias

Tests.—The distinguishing tests for Dill Oil are the strong characteristic odour, the specific gravity, which should be 0.905 to 0.920, the optical rotation, which should be from $+70^{\circ}$ to $+80^{\circ}$, in a tube of 100 mm length. It should yield a clear solution in from 6 to 8 parts of Alcohol (80 p.c.). Not less than 15 p.c. should distil below 185°C (365°F), and not less than 40 p.c. above 220°C (428°F). The abstraction of Carvone from the oil is indicated by a decrease in the specific gravity, and a diminution in the amount distilling above 220°C (428°F).

Not Official

ANILINE

$\text{C}_6\text{H}_5\text{N}$, eq. 92.40

* An oily liquid, colourless when freshly distilled, but very prone to become yellow or brown on exposure to air.

It should be kept in well stoppered bottles of a dark amber tint and protected as far as possible from the air.

Solubility—1 in 27 of Water, 5 in 4 of Alcohol (60 p.c.), mixes in all proportions with Alcohol (90 p.c.), Ether and Glycerin.

As a vehicle for dissolving Cocaine, for the production of local anaesthesia of the ear, 10 to 15 minims of a 5 p.c. solution of Cocaine made by dissolving Cocaine Hydrochloride 5 in dilute Alcohol 50, Aniline Oil 50—*L* '00, 1 1125, '01, 1 698.

Dose—Not more than 7 minims = 0.5 c.c.—*L* '00, 1 1127.

Several cases of poisoning from boots to which a black material containing an Aniline dye had recently been applied—*L* '02, 1 463.

Tests—It should possess a specific gravity of 1.020 to 1.026, and a boiling point of 183° to 184°C (361.4° to 363.2°F). An aqueous solution of Aniline treated with a solution of Chlorinated Lime yields a dirty violet blue coloration changing to Indigo blue on the addition of Ammonia Solution. A few drops of Aniline warmed with an alcoholic solution of Potassium or Sodium Hydroxide and a drop or two of Chloroform evolve the characteristic, disagreeable, poisonous odour of Phenyl Isonitrile.

ANISI FRUCTUS.

ANISE FRUIT

Fr, ANIS VLRT, GER, ANISSAMEL, ITAL, ANICF, SPAN, ANIS

The dried ripe Fruit of *Pimpinella Anisum*, L.

Medicinal Properties—Stomachic and aromatic, carminative, slightly expectorant, used to relieve flatulence, and to diminish the griping of purgative medicines.

Official Preparations—Aqua Anisi and Oleum Anisi.

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex (Anis Comun), Norw, Port, Russ, Span, Swed, Swiss and U.S.

Descriptive Notes.—Anise fruit occurs in commerce in several varieties, differing considerably in size and colour, and in degree of freedom from impurity. It possesses an aromatic and characteristic

odour, and has a sweet taste. The official drug is limited to such varieties as are about $\frac{1}{2}$ inch (5 mm) long, and $\frac{1}{16}$ inch (2 mm) broad, and therefore includes the Maltese, Alicante and German varieties. Anise fruit is greyish-brown, *BP* (also greyish or greenish-grey, *USP*), ovate, stalked, with the two carpels united, and rough with minute 1-celled hairs, a transverse section exhibits numerous vittæ. The German variety is brownish and that of Alicante greenish, South Russian Anise is similar in size to *Conium* fruit. *Conium* has sometimes occurred mixed with Anise fruit in Italy. Its presence may be detected by the mouse-like odour developed when rubbed with a few drops of *Liquor Potassæ*. It differs from Anise in being without hairs, and having the ridges distinctly crenate, and the flat surface deeply grooved as seen in transverse section, and in being without vittæ in the mature fruit.

North Russian Anise is small and dark green or brownish, and it is used as a cheap source of the essential oil. The Syrian and Chilean varieties are usually very inferior and mixed with more or less foreign matter, and are consequently reserved for veterinary medicine. The varieties richest in essential oil are the Italian, Spanish and South Russian. The residue after distillation is valued as an ingredient for cattle foods. Under the microscope the distinguishing features are the simple, thick-walled, short, erect, straight or slightly curved hairs, with a minutely warty surface, and the striated surface of the flattened polygonal cells of the outer epidermis.

AQUA ANISI. ANISE WATER

Anise Fruit, 1, Water, 20, distil, 10 (1 in 10)

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 c.c.

Foreign Pharmacopœias.—Official in Belg., from the Spirit, Fr., Jap., Port and Span., from Fruits, and U.S. from Oil. Not in the others.

OLEUM ANISI. OIL OF ANISE

Fr., Essence d'Anis, Ger., Anethol, Ital., Essenza di Anice, Span., Esencia de Anis.

At temperatures above 15°C (59°F) it is a colourless or pale yellow refractive liquid, with a pleasant aromatic odour and very sweet taste, below 15°C (59°F) it becomes a white crystalline solid. It is obtained by distillation from the fruits of the official variety, or from the fruit of *Illicium Verum* or Star Anise.

It should be kept in amber-coloured well-stoppered glass bottles.

Solubility.—1 of Pimpinella Oil in 3 of Alcohol (90 p.c.); 1 of Illicium Oil in 4 of Alcohol (90 p.c.), a slight rise in temperature greatly increases the solubility in Alcohol (90 p.c.), both oils dissolve in all proportions of Absolute Alcohol, 1 of Pimpinella Oil in 200 of Alcohol (60 p.c.), at which point the Illicium Oil is distinctly turbid.

These variations in solubility seem to arise from the presence in the Illicium Oil of a small proportion of a much less soluble Oil, which is absent in the Pimpinella.

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Prescribing Notes—*May be taken on Sugar Before dispensing an oil which has become partly solidified, it should be completely liquefied by warming, and well mixed by shaking*

Official Preparation—Spiritus Anisi Contained in Tinctura Camphoræ Comp and Tinctura Opi Ammoniata

Not Official—Aniseed Cordial, Elixir Anisi, Essentia Anisi, Tinctura Anisi, Anisic Acid, Sodium Anisate and Anethol

Foreign Pharmacopœias—The following are from *Pimpinella* Dan, Dutch, Fr, Ital, Norw and Russ, sp gr 0 980 to 0 990, Hung, sp gr 0 978 to 0 984, Port, sp gr 0 977 to 0 983, Mex, sp gr 0 982, Span, sp gr 0 984 to 0 986, Swiss, sp gr 0 984 to 0 994, U S, sp gr 0 975 to 0 985 at 25° C (77° F) The following permit the use of both kinds Mex, Port and U S See also Anethol

Tests.—The distinguishing tests for Anise Oil are its distinctive odour, its sweet taste, the specific gravity, which at 20° C (68° F) should be from 0 980 to 0 990, its optical rotation, which should be -1° to -2° in a tube of 100 mm length, the melting point, which should not be below 15° C (59° F), the solidifying point, which should not be below 15° C (59° F)

The *BP* gives the sp gr at 20° C (68° F) as 0 975 to 0 990, the *PG* at 25° C (77° F) as 0 984 to 0 986, the *USP* at 25° C (77° F) as 0 975 to 0 988 The melting point given in the *BP* is from 10° to 15° C (50° to 59° F), the *USP* states that it should not be below 15° C (59° F) and gives the method outlined below for its determination

80 to 90 pc of the oil should distil between 225° and 235° C (437° to 455° F), indicating a due percentage of Anethol Neither *BP* nor *USP* make any reference to either the boiling point or to fractionation

The *Pimpinella* Oil is readily distinguished from that of *Star Anise* by giving a deep blue colour on the addition of saturated solution of Hydrochloric Acid gas in Alcohol, but Schimmel and Co state that the two varieties can only be distinguished by the odour and taste, and that the reaction with alcoholic Hydrochloric Acid does not give reliable results The balance of opinion seems to be that it does afford a means of distinction

The more generally occurring sophistications are Fennel Oil, or its stearoptene, volatile oils containing Phenols, Alcohol, the fluid portion remaining after the extraction of the Anethol, and Petroleum Fennel Oil or its stearoptene are detected by the optical rotation of the sample, both these adulterants being dextrogyrate, volatile oils containing Phenols are detected by a blue or brownish colour produced by the addition of Ferric Chloride Test solution to an alcoholic solution of the oil, Alcohol by the decrease in volume when the oil is shaken in a graduated measure with Water, the fluid portion remaining after the extraction of Anethol, by the alteration in the melting and solidifying point, and Petroleum by the Alcohol solubility—the pure oil dissolves 1 in 3 of Alcohol (90 pc), an oil containing 5 pc of Petroleum will not dissolve 1 in 10 The oil undergoes oxidation by exposure to air, and its characters are greatly changed. Rise of specific gravity and lowering of the melting point are the principal indications as to the extent of oxidation.

Melting Point.—In the *USP* the following method is given to determine the congealing point, which should be below 15°C (59°F). Transfer about 10 c.c. of the Oil to a test-tube placed in Water cooled with Ice, insert a thermometer at once into the Oil and allow it to remain undisturbed until its temperature has fallen to about 6°C (42.8°F). Induce crystallisation either by rubbing the inner wall of the test-tube with the thermometer or by the addition of a particle of Solid Anethol, remove the test-tube from the bath, and stir constantly during the solidification of the Oil. The highest temperature reached during the crystallisation is regarded as the congealing point.

Preparation.

SPIRITUS ANISI. SPIRIT OF ANISE

Oil of Anise, 1, Alcohol (90 p.c.), *q.s.* to make 10 (1 in 10)

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

Half the strength of *BP* '85.

Foreign Pharmacopœias.—*Fr.* Oil in 100, *Fr.* 1 Oil in 50, *US*, Spirit, 1 Oil in 4, *Span.* has Alcohol de Anis Amoniacal, Oil 1, Liquid Ammonia 5, Alcohol (95 p.c.) 24. All by weight, except *US*. Not in the others.

Not Official.

ELIXIR ANISI. *Syn.* ANISEED CORDIAL.—Anethol, 3.5, Oil of Fennel, 0.5, Spirit of Bitter Almond (*USP*), 12, Alcohol (95 p.c.), 240, Syrup (*USP*), 625, Water, 125, Purified Talc (*USP*), 15. Average Dose for infants, 1 c.c. (15 minims).—*USNF*

Anethol, 0.35, Oil of Fennel, 0.05, Spirit of Bitter Almond, 1.25; Alcohol (90 p.c.), 24, Syrup, 62.50, Magnesium Carbonate, 1.50, Distilled Water, *q.s.* to produce 100.—*BP C*

ESSENTIA ANISI—Oil of Anise, 1, Rectified Spirit, 4.—*BP* 1885
This has been incorporated in the *BP C*

TINCTURA ANISI—(*Ital*, *Mex* and *Russ*)—Anise Fruit, 1, Alcohol, 5

ANISIC ACID ($\text{H}_2\text{C}_6\text{H}_3\text{O}$)—It occurs in colourless, shining acicular crystals obtained by the oxidation of Oil of Anise or Anethol

Solubility—Almost insoluble in cold Water, 1 in 700 boiling Water, 1 in 36 of Alcohol (90 p.c.), 1 in 50 of Ether

SODIUM ANISATE—In rhombic crystals, or a crystalline powder, frequently efflorescent, with a slight aromatic odour

Solubility—1 in 5 of Water, 1 in 24 of Alcohol (90 p.c.)

Anisic Acid and its Sodium salt have been stated to possess antiseptic and antipyretic properties, similar to Salicylic Acid

Dose—5 to 15 grains = 0.32 to 1 gramme

ANETHOL ($\text{C}_{10}\text{H}_{12}\text{O}$)—The Stereoptene separated from either of the Anise Oils. It is said to have a finer flavour than the Oil, being free from the acidity pertaining to the non-freezing portion of the Oil. A white crystalline mass possessing a strong characteristic Anise-like odour and sweetish taste. When melted it forms a highly refractive, almost colourless liquid

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Ger., Jap. and Swed. Not in the others

Tests.—It has at 25°C (77°F) a sp. gr. of 0.954 to 0.966, a melting point of 22.5° to 23°C (72.5° to 73.4°F), and a boiling point of 232° to 234°C (449.6° to 453.2°F). It should form a clear solution in 3 parts of Alcohol (90 p.c.).

ANTHEMIDIS FLORES.

CHAMOMILE FLOWERS

FR, CAMOMILLE ROMAINE, GTR, ROMISCHE KAMILLE, ITAL, CAMOMILLA ROMANA, SPAN, MANZANILLA ROMANA

The dried expanded Flower heads of the common or Roman Chamomile, *Anthemis nobilis*, L., collected from cultivated plants

Medicinal Properties—Stomachic, tonic and carminative In large doses, emetic Useful in atonic dys-pepsia, externally it is employed as a fomentation for bruises and contusions

Prescribing Notes—*The Extract or Oil is frequently added to Rhubarb and aperient medicines as a corrective A little Soap added in the case of the Oil, makes a good pill mass*

Official Preparations—Extractum Anthemidis, and Oleum Anthemidis The Oil is contained in the Extract

Not Official—Aqua Anthemidis, Decoctum Anthemidis et Papaveris, Oleum Chamomillæ Infusum, Infusum Anthemidis, Infusum Anthemidis Concentratum, Liquor Anthemidis et Papaveris, and Tinctura Anthemidis

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Ital, Jap, Mex, Port, Span, Swiss and U S Not in the others Also *Matricaria* in Austr, Dan, Dutch, Ger, Hung, Ital, Jap, Mex, Norw, Russ, Span (*Manzanilla ordinaria*), Swed and Swiss

Descriptive Notes—The drug is met with in commerce chiefly in two forms, known respectively as English and exotic The official description as well as that of the *USP* applies to the former The flower-heads are hemispherical, 10 to 20 mm ($\frac{1}{2}$ to $\frac{3}{4}$ inch) in diameter, and nearly white in colour The florets are ligulate, suddenly tapering in the lower half and about 1 cm ($\frac{1}{2}$ inch) long, a few tubular florets usually remaining in the centre There is no pappus, and the solid conical receptacle is covered with narrow membranous bracts or paleæ, visible when the florets are removed. The involucre scales are also membranous and obtuse, with a green central nerve The exotic Chamomile flowers, chiefly imported from Belgium and France, differ in being rather larger and in the florets being broader and all ligulate In Scotland the single wild Chamomile flowers are often sold under the name of Scotch Chamomiles These differ in having only one row of ligulate florets, all the centre consisting of yellow tubular florets Scotch Chamomiles are sometimes preferred, on the supposition that the central contain more oil than the ligulate florets, but the single Chamomiles are excluded from use in dispensing by the *BP* description

The Chamomile flowers (Flores Chamomillæ) official in the *PG*, are the single flower-heads of *Matricaria Chamomilla*, L These are much smaller than those of *Anthemis nobilis*, averaging only $\frac{1}{2}$ inch (5 mm) in diameter, and have a portion of the fruit-stalk attached to them The receptacle is nearly conical, but hollow, and has no paleæ The odour resembles that of the *BP* Chamomile, but is weaker

If Chamomile flowers have a brownish tint they have either been damaged by rain, or not carefully dried, or have become brownish by

keeping. In either case they must be regarded as of inferior quality and unfit for medicinal purposes. The taste of Chamomile flowers is bitter and strongly aromatic.

The double flowers of *Chrysanthemum Parthenium*, Bernh., which have occasionally been found mixed with Chamomile flowers have a slightly convex, not conical, solid receptacle, and the few paleæ sometimes present are lanceolate and acute.

Preparations

EXTRACTUM ANTHEMIDIS. EXTRACT OF CHAMOMILE.

An aqueous extract of the Flowers treated by decoction, to which Oil of Chamomile is added just before completion of the evaporation, in the proportion of 15 minims of Oil to each pound of the Flowers employed.

The double Flowers yield about 30 p.c. of Extract.

Dose.—2 to 8 grains = 0.13 to 0.52 gramme.

Foreign Pharmacopœias—Official in Ital., from both, Dan. and Swed., from Matricaria, Mex. (Extracto de Manzanilla). Not in the others.

OLEUM ANTHEMIDIS. OIL OF CHAMOMILE.

A pale blue oily liquid, of an aromatic characteristic odour and slightly bitter taste, distilled from the dried Flowers of *Anthemis nobilis*. On exposure to light and air the Oil acquires a greenish or yellowish-brown colour, and should therefore be kept in well-stoppered glass bottles of a dark amber colour, and exposed as little as possible.

Solubility—Scarcely in Water, 10 in 3 of Alcohol (90 p.c.)

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Foreign Pharmacopœias—Official in Belg., from Anthemis, Fr., from Anthemis, Ital. and Swiss, from Matricaria. Not in the others.

Tests.—Oil of Chamomile has a specific gravity of about 0.910. The optical rotation should be between $+1^{\circ}$ and $+3^{\circ}$ in a tube of 100 mm. length. The optical rotation is not given in the B.P.

Not Official.

AQUA ANTHEMIDIS—Flowers 1, Water 20, distil 10. (1 in 10)

Foreign Pharmacopœias—Official in Austr., 1 in 10; Belg., 8 of the Spirit in 100, Port., 1 in 4. Ital., Matricaria, 1 in 2, Anthemis, 1 in 2 $\frac{1}{2}$. All distilled. Port., from Anthemis, Austr., from Matricaria, Ital., from both. Not in the others.

Oleum Chamomillæ Infusum—Chamomile Flowers, 1, Olive Oil, 10; digest in a water-bath for 2 hours, strain, press and filter.

Foreign Pharmacopœias—Official in Fr. and Port., 1 in 10; Span., 1 in 8, from Anthemis. Ital., 1 in 4, from Anthemis and from Matricaria. Belg. (Oleum Chamomillæ Camphoratum), 1 of Oil with 999 of Camphorated Oil. Span. (Aceite de Manzanilla Alcanforado), 1 of Camphor with 9 of Oil Cham. Infus.

DECOCTUM ANTHEMIDIS ET PAPAVERIS—Chamomile Flowers, 10; Poppy Capsules, bruised, 5, Distilled Water, q.s. to produce 100.—B.P.C.

INFUSUM ANTHEMIDIS—Chamomile Flowers, 1, Boiling Distilled Water, 20 Infuse 15 minutes and strain—*B P* 1885
This has been incorporated in the *B P C*

INFUSUM ANTHEMIDIS CONCENTRATUM—Chamomile Flowers, in powder, 40, Oil of Chamomile, 0.2, Alcohol (20 p c), *q s* to make 100, mix the Oil of Chamomile thoroughly with the powder and submit the latter to re-percolation

Dose—As a stomachic, 1 to 4 fl drim, as an emetic, 5 to 10 fl drim—*Farr* and *Wright*, *P J* '06, 1 165 and '07, 1 621, *C D* '06, 1 252, *Y B P* 1907, 249
This appears in the *B P C*

LIQUOR ANTHEMIDIS ET PAPAVERIS—Concentrated Infusion of Chamomiles and Liquid Extract of Poppies, equal volumes One or two tea spoonfuls in half a pint of boiling Water, for use as a fomentation

TINCTURA ANTHEMIDIS—Single Chamomiles, carefully dried, 1, sufficient Alcohol (90 p c) to percolate, 8, or an equivalent quantity of fresh Flowers (about 3), and macerate with 8 of Alcohol (90 p c) for 7 days, and press.

The moisture in the fresh flowers reduces the strength of the spirit so that less resin is dissolved, and the tincture is consequently less bitter

Foreign Pharmacopœias—Official in Austr, 1 in 5, from Matricaria, Ital and Mex, 1 in 5, both varieties, Belg, Chamomillæ Spiritus, 1 of the Oil in 100

ANTIMONII OXIDUM.

ANTIMONIOUS OXIDE

Sb_4O_6 , eq 571.28

A white, more or less crystalline, powder

When a solution of Antimonious Chloride is poured into Water Antimony Oxichloride is precipitated, and this, when in turn decomposed with Sodium Carbonate, yields Antimonious Oxide

Solubility.—Insoluble in Water, Alcohol, and Nitric Acid, readily dissolved by Hydrochloric Acid and warm solution of Tartaric Acid

Medicinal Properties—Similar to but less active than the Tartrate because less soluble

Dose—1 to 2 grains = 0.06 to 0.13 gramme

Prescribing Notes—The *Pulvis Antimonialis* is generally given in the form of powders, pills or cachets

Official Preparation.—*Pulvis Antimonialis* Used in the preparation of Antimonium Tartaratum

Foreign Pharmacopœias—Official in Mex (*Oxido Antimonioso Precipitado*), Norw (*Oxydum Stibicum*), Port and Span Not in the others

Tests—The distinguishing tests for Antimony Oxide are that its slightly acidified solution yields with Hydrogen Sulphide Solution an orange-coloured precipitate soluble in Potassium Hydroxide Solution and in Ammonium Hydrosulphide Solution, insoluble in Ammonium Carbonate Solution, a cold white porcelain vessel allowed to impinge upon the upper flame produced on igniting the gas yielded by the

interaction of Zinc, Hydrochloric Acid and a solution of the Oxide, acquires a dark metallic-looking stain, which is unaffected by Calcium or Sodium Hypochlorite Solution, the production of a metallic-looking coating on Copper when a piece of Copper foil is boiled with a solution of the Oxide containing free Hydrochloric Acid, the formation of a white amorphous but no crystalline sublimate when this coating is volatilised in a tube, the formation of a black deposit of Antimony on the Platinum, when a Zinc rod is allowed to rest on a piece of Platinum foil in an acidified solution of the Oxide.

It is officially required to indicate 99.98 p.c. of pure Antimonious Oxide, as indicated by titration with Deci-normal Volumetric Iodine Solution. The Oxide is brought into solution by means of Acid Potassium Tartrate or Tartaric Acid and an alkaline reaction throughout the titration ensured by the addition of Sodium Bicarbonate, this being necessary to neutralise the Hydriodic Acid set free during the reaction, Hydriodic Acid otherwise acting as a reducing agent. A weighed quantity of 0.5 gramme dissolved in twice the volume of Acid Potassium Tartrate then mixed with 3 grammes of Sodium Bicarbonate and cooled should decolorise 70 c.c. of the Volumetric Solution of Iodine.

The more generally occurring impurities are Antimonic compounds, Arsenic, Calcium, Copper, Iron, Lead, Chlorides and Sulphates. It is officially required to dissolve completely in an excess of Acid Potassium Tartrate. The extent to which it is true depends upon the age of the specimen, Antimonious Oxide undergoes oxidation on exposure to air, the resulting Antimonic compounds being insoluble. The amount of residue therefore forms a criterion of the proportion of the latter. The remaining impurities are grouped together in the *BP* under the elastic expression, 'it should yield no characteristic reaction with the tests for,' etc. The solution obtained by dissolving the Oxide in Hydrochloric Acid should yield no reaction with the Bettendorf's test, indicating the absence of Arsenic. When dissolved in Sodium Hydroxide Solution (15 p.c. w/w) it should yield on the addition of Hydrogen Sulphide Solution no dark coloration, indicating the absence of Copper, Lead and Iron, a white precipitate would indicate the presence of Zinc. Another portion of the sample dissolved in Sodium Hydroxide Solution (15 p.c. w/w) when supersaturated with Nitric Acid should yield only a faint turbidity on the addition of either Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than a trace of Chlorides and Sulphates.

Preparation

PULVIS ANTIMONIALIS. ANTIMONIAL POWDER.

Antimonious Oxide, 1, Calcium Phosphate, 2.

Dose.—3 to 6 grains = 0.2 to 0.4 gramme

Foreign Pharmacopœias — Official in Mex., Antimonious Oxide 1, Calcium Phosphate 2, Port., Antimonious Oxide 85, Calcium Phosphate 65, and in the others.

ANTIMONIUM NIGRUM PURIFICATUM.

ANTIMONIOUS SULPHIDE

FR, TRISULFURE D'ANTIMOINE, GER, GEREINIGTES SCHWEFELANTIMON;
ITAL, TRISOLFURO DI ANTIMONIO, SPAN, SOLFURO DE ANTIMONIO PURIFICADA.

A heavy, odourous, greyish-black crystalline powder, consisting of native Antimonious Sulphide Sb_2S_3 , eq 333 46, separated from siliceous material and from arsenical compounds

Official Preparation —Used to prepare Antimonium Sulphuratum

Foreign Pharmacopœias —Official in Austr, Belg, Dan, Fr, Ger, Hung, Ital (Ciudo and Depurato), Mex, Poit, Span (Sulfuro de Antimonio Purificada), Swed and Swiss Not in the others

Tests —Antimonious Sulphide yields the tests distinctive of Antimony given under Antimonious Oxide, it moreover evolves on boiling with Hydrochloric Acid the characteristic disagreeable odour of Hydrogen Sulphide, and when fused with a mixture of Sodium Carbonate and sufficient Potassium Nitrate to oxidise it, yields a mass which, when dissolved in Water and filtered, affords with Barium Chloride Solution a heavy white precipitate insoluble in Hydrochloric or Nitric Acid, or in a mixture of both

The more generally occurring impurities are Arsenic and siliceous matter The Pharmacopœia stipulates that it should not yield more than slight characteristic reactions with the Arsenic tests The Belgian Pharmacopœia is more precise, and requires that 1 gramme, shaken with 10 c c of Hydrochloric Acid until disengagement of gas ceases, mixed with 5 c c of Water, and cautiously heated on a water-bath with some crystals of Potassium Chlorate, the mixture, when freed from Chlorine, filtered through an asbestos plug and evaporated to 3 c c, shall yield no brown coloration in one hour, when mixed with an equal volume of Stannous Chloride Solution The *P G* and Austri Ph do not allow more than 1 p c of residue insoluble in concentrated Hydrochloric Acid

Hydrochloric Acid —2 grammes of the powdered salt are dissolved on warming gently with 20 c c of Hydrochloric Acid, and then, on boiling for 1 hour, should leave a residue amounting to not more than 0 02 gramme, *P G*

ANTIMONIUM SULPHURATUM. SULPHURATED ANTIMONY *N O Syn* —KREMES MINERAL STIBIUM SULFURATUM AURANTIACUM

FR, PENTASULFURE D'ANTIMOINE, GER, GOLDSCHWEFEL, SPAN, AZUFRE DORADO DE ANTIMONIO

An orange-red, odourless, tasteless powder, which consists of a mixture of Antimony Penta- and Tri-sulphides and -oxides, and containing also some free Sulphur It should be preserved from the light

It is officially stated to be prepared by boiling 10 parts each of Antimonious Sulphide and Sublimed Sulphur for two hours with a solution of 5 parts of commercial Sodium Hydroxide in 100 parts of Distilled Water, stirring frequently, the volume of the liquid is maintained by the occasional addition of Distilled Water Whilst still hot, add 180 parts of boiling Distilled Water, strain through calico,

and add gradually diluted Sulphuric Acid in slight excess to the strained liquid. Collect the precipitate on calico, wash with Distilled Water till the washings are free from Sulphates, and dry at a temperature not higher than 100° C (212° F)

Solubility.—Insoluble in Water, dissolves readily in Sodium Hydroxide solution also in hot Hydrochloric Acid evolving Hydrogen Sulphide

Medicinal Properties.—Alterative, diaphoretic, and emetic, powerfully depressant, uncertain in action from its slight solubility, depending on the acidity of the stomach. Usually prescribed with Calomel and Guaiacum, as in *Pilula Hydrargyri Subchloridi Composita*, as a cholagogue in gout, for secondary syphilis and its cutaneous eruptions, or with Henbane or Hemlock in chronic rheumatism

Objection taken (*L* '05, 1 1610) to the use of red Antimony Sulphide in the composition of red India-rubber. The investigation arose from a case of appendicitis where a large quantity of liquid was in the habit of being drunk from bottles fitted with stoppers having a red rubber ring round them

It is pointed out (*L* '05, 1 1786) that this Sulphide is soluble with difficulty except in hot strong acids or strong alkali, and a series of tests carried out with a number of the more generally occurring beverages, at which they possessed no solvent action as regards Antimony when left in contact with red rubber rings containing 16.83 p.c. of that substance

Dose.—1 to 2 grains = 0.06 to 0.13 gramme

Official Preparation.—Contained in *Pilula Hydrargyri Subchloridi Composita*

Foreign Pharmacopœias.—Official in Austr, Belg, Hung and Swiss (*Stibium Sulfuratum Aurantiacum*), Dan and Dutch (*Sulfidum Stibicum*), Fr (*Pentasulfure d'Antimoine*), Ger, Jap and Russ (*Stibium Sulfuratum Aurantiacum*), Mex (*Sulfuro Antimonico*), Norw (*Sulfuretum Stibicum*), Port (*Enxofre Dourado de Antimonio*), Span (*Azufre Dorado de Antimonio*), Swed (*Kermes Mineralis*). Swiss has also *Stibium Sulfuratum Rubrum*

Tests.—Sulphurated Antimony yields the tests distinctive of Antimony appearing under Antimonious Oxide, when treated with Hydrochloric Acid it evolves a disagreeable characteristic odour of Hydrogen Sulphide, and a separation of Sulphur occurs; and when fused with Potassium or Sodium Carbonate and sufficient Potassium Nitrate to effect oxidation, the product, when dissolved in Water and filtered, yields a solution giving with Barium Chloride Solution a dense white precipitate insoluble in concentrated Hydrochloric Acid or in Nitric Acid, or in a mixture of both

The official gravimetric test requires that when 3 grammes of the Sulphide are fully oxidised with Nitric Acid, 2 grammes of Antimonious Oxide should be produced. It has been pointed out (*Y.B.P.* '07, 473) that it will not yield this amount of residue by this test, and it has been suggested that it should be modified so as to read, 3 grammes moistened with dilute Nitric Acid, with successive portions of fuming Nitric Acid until red fumes cease to be evolved, the excess of Water then evaporated off, and carefully heated to redness, to expel Sulphuric

Acid, should leave a whitish residue weighing not less than 1.6 and not more than 1.8 grammes. It is shown that a sample containing as much as 30 per cent of anhydrous Sodium Sulphate yielded a figure for residue very close to that obtained from a genuine and carefully prepared sample.

The more generally occurring impurities are Arsenic, siliceous matter, Chlorides and Sulphates. The *B P* employs the ordinary tests for Arsenic. The *P G* uses saturated aqueous Ammonium Carbonate Solution for the extraction of the Arsenic, and states that when this solution is supersaturated with Hydrochloric Acid no yellow flocculent precipitate should be thrown down in six hours. The Belgian Pharmacopœia gives an Arsenic test similar to that recorded under Antimonious Sulphide. The tests described in small type below, under the heading of Silver Nitrate and Barium Nitrate, serve to detect Chlorides and Sulphates if present.

Ammonium Carbonate Solution—Allow a mixture of 0.5 gramme of Sulphurated Antimony with 5 cc of an aqueous Solution of Ammonium Carbonate saturated at ordinary temperatures, to stand for 2 minutes with occasional agitation at a temperature of 50° to 60° C (122° to 140° F). In the solution so obtained after filtration, and saturation with Hydrochloric Acid, a yellow flocculent precipitate should not be thrown down within six hours, *P G*.

Silver Nitrate—1 gramme Sulphurated Antimony when agitated with 20 cc of Water and filtered, gives a filtrate which on the addition of Silver Nitrate T.S. should become only faintly opalescent, but should not become brown, *P G*.

Barium Nitrate—The filtrate obtained as above should not immediately become cloudy on the addition of T.S. of Barium Nitrate, *P G*.

Not Official

LIQUOR ANTIMONII CHLORIDI—A yellowish red liquid, sp gr about 1.47. A powerful escharotic.

ANTIMONIUM TARTARATUM.

TARTARATED ANTIMONY

B P Syn—POTASSIO TARTRATE OF ANTIMONY, TARTAR ÉMÉTIQ.

FR, ANTIMONIO-TARTRATE ACIDE DE POTASSIUM, GER, BREI CHWEINSLEIM
ITAL, TARTARO EMETICO, SPAN, TARTRATO ANTIMONICO POTASICO

$[K(SbO)C_4H_4O_6]_3H_2O$, eq 659.14.

N O Syn—TARTARUS STIBIATUS

Colourless, odourless, transparent rhombic crystals, or as a heavy white powder. Taste at first sweet, then nauseous and metallic.

It should be preserved in well-stoppered bottles of a dark amber colour.

Solubility.—1 in 17 of Water (slowly), 1 in 2 of boiling Water, sparingly soluble in Alcohol (60 p.c.), insoluble in Alcohol (90 p.c.)

Medicinal Properties—Diaphoretic, expectorant, alterative, emetic, circulatory and nervous depressant. Useful in the head

symptoms of acute febrile diseases and in delirium tremens, contra-indicated in asthenic cases, alterative in chronic skin affections and in gout

As a diaphoretic and expectorant, it is given with great effect in the treatment of acute pneumonia, bronchitis and croup

Externally, in the form of ointment, it acts as a powerful counter-irritant, producing a pustular eruption

Too purgative and depressant for use in lowering blood pressure B M J '06, 11 1451

A review of post-mortem examinations, after death from the administration of tartar emetic—B M J '03, 1 873

Dose.—As a diaphoretic, $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.003 to 0.008 gramme; as an emetic, 1 to 2 grains = 0.06 to 0.13 gramme

Ph Ger maximum single dose, 0.2 gramme, maximum daily dose, 0.6 gramme

Prescribing Notes.—*Best prescribed in aqueous solution or as the Vinum. In pill, well triturated with Milk Sugar and Diluted Glucose q s*

Incompatibles.—Tannic Acid, the Alkalis and their Carbonates, and Lead salts, Astringent infusions, as Cinchona, Rhubarb, etc

Official Preparation.—Vinum Antimoniale

Not Official.—Unguentum Antimonii Tartarati.

Antidotes.—Stomach-tube or emetics, Tannic Acid, Catechu, vegetable astringents, Tea or Coffee, stimulants if much collapse

Foreign Pharmacopœias.—Official in Austr (Stibium Kalio-Tartaricum), Belg (Tartarus Stibiatus), Dan, Nor and Swed (Tartarus Stibico-Kalicus), Dutch (Tartras Kalico-Stibicus), Fr (Antimoniotartrate Acide de Potassium), Ger and Swiss (Tartarus Stibiatus), Hung (Kalium Stibio-Tartaricum), Ital (Tartrato di Antimonio e di Potassio), Mex (Tartrato de Potassio y antimonio), Port (Tartrato de Potassa e de Antimonio), Jap. and Russ (Stibio-Kalium Tartaricum), Span (Tartrato Antimonico Potassico), US (Antimonii et Potassii Tartras)

Tests.—Tartarated Antimony should answer the tests distinctive of Antimony given under Antimonious Oxide, after separation of the Antimony it should also give a yellow crystalline precipitate with Platinic Chloride Solution, and a residue of Platinum and Potassium Chloride when this precipitate is ignited, when moistened with Hydrochloric Acid and introduced into a Bunsen flame it should communicate a violet coloration readily distinguished when viewed through a blue glass, it should give a white precipitate soluble in moderately concentrated Potassium Hydroxide Solution, when an aqueous solution is tested with Calcium Chloride Solution; with Silver Nitrate Solution a white precipitate soluble in Ammonia Solution, the ammoniacal solution yielding a precipitate of metallic Silver when the solution is boiled, the precipitate is also soluble in Nitric Acid, a purple or violet colour should be produced when to its solution acidulated with Acetic Acid, a drop or two of Ferrous Sulphate Solution is added, followed by a few drops of Hydrogen Sulphide Solution, and then an excess of Potassium Hydroxide

Solution The *BP* states that an aqueous solution is not precipitated by Gallic Acid, but this is contrary to general experience. A note in reference to this will be found under Gallic Acid. It is officially required to indicate not less than 99.19 p.c. and not more than 100.02 p.c. of the pure salt as ascertained by titration with Decinormal Volumetric Iodine Solution, an excess of Sodium Bicarbonate being maintained throughout the operation in order to neutralise the Hydriodic Acid produced during the reaction. The Sodium Bicarbonate must be added not long before the titration or the Antimony will be precipitated. A weighed quantity of 1 gramme when mixed with about 3 times its weight of Sodium Bicarbonate and dissolved in Water should require not less than 60.2 nor more than 60.7 c.c. of the Volumetric Solution.

The *USP* Volumetric test indicates not less than 99.5 p.c. of the pure salt, the *PG* indicates 99.6 p.c., both processes are compared in small type below under the heading Volumetric Determination.

It is officially required that 1.66 grammes should slowly dissolve without leaving a residue, in 25 c.c. of Water at 15.5° C (60° F).

The more generally occurring impurities are Ammonium salts, Arsenic, Calcium, Copper, Iron, Lead, Sodium, Chlorides, Sulphates and Potassium Acid Tartrate. In the *BP* these are grouped collectively. The more important are Arsenic, Copper and Lead, Iron and Potassium Acid Tartrate. Both *USP* and *PG* adopt the Bettendorf's test for Arsenic. Heavy metals, *e.g.*, Copper, Lead and Iron may be detected by adding to an aqueous solution sufficient Potassium or Sodium Hydroxide Solution to redissolve the precipitate at first formed and then passing Hydrogen Sulphide into the liquid. Chlorides and Sulphates respond to the usual tests, Potassium Acid Tartrate may be detected by the effervescence produced on adding Sodium Bicarbonate Solution. The *PG* includes a test for Arsenic, but for no other impurity. The 1 in 100 aqueous solution acidified with Acetic Acid should be unaffected by Ammonium Oxalate Solution, indicating the absence of Calcium. The *USP* includes a separate test for Iron, which is given in small type under the heading of Potassium Ferrocyanide Solution.

Residue—On heating to 110° C (230° F) it loses its Water of crystallisation (2.71 p.c.) and at a red heat chars, emitting an odour resembling that of burning Sugar, and leaving a black residue with an alkaline reaction, *USP*, it chars on heating, *PG*.

Potassium Ferrocyanide Solution—An aqueous 1 in 100 solution acidulated with Acetic Acid should be unaffected by T.S. of Potassium Ferrocyanide, *USP*.

Stannous Chloride Solution—A mixture of 1 gramme Tartarated Antimony and 3 c.c. Stannous Chloride Solution should not assume a dark colour in the course of an hour, *PG*, 2 grammes dissolved in 5 c.c. Hydrochloric Acid should not respond to Bettendorf's test for Arsenic, *USP*.

Hydrogen Sulphide Solution—If sufficient solution of Sodium Hydroxide be added to an aqueous solution (1-20) of Tartarated Antimony to redissolve the precipitate first formed, and then an equal volume of freshly prepared Hydrogen Sulphide Solution added, no coloration should be noticeable after standing in a warm place for half an hour, when viewed by reflected light, holding it against a white surface, indicating absence of heavy metals, *USP*.

Sodium Bicarbonate or Carbonate—No effervescence should occur with Solution of Sodium Bicarbonate, *B P*, Solution of Sodium Carbonate, *U S P*. Absence of Acid Potassium Tartrate

Volumetric Determination—1 gramme is dissolved in sufficient Water to measure 100 cc, then 33 cc of this solution should, after the addition of 20 cc of a cold saturated aqueous solution of Sodium Bicarbonate and a little Starch T S, require not less than 19.9 cc of Tenth-normal Iodine V S. to produce a permanent blue colour (each cc of the pure salt). Titration should begin immediately after the addition of the Sodium Bicarbonate Solution, *U S P*. A solution of 0.2 gramme of Tartarated Antimony and 0.2 gramme of Tartaric Acid in 100 cc of Water should after the addition of 2 grammes of Sodium Bicarbonate and a few drops of Starch Solution, require 12 cc of Tenth-normal Solution of Iodine to produce a blue colour, *P. G*

Preparation.

VINUM ANTIMONIALE. ANTIMONIAL WINE.

Tartarated Antimony, 40 grains, boiling Distilled Water, 1 fl oz, Sherry, *q s* to yield 20 fl oz

Boiling Water is added to dissolve the Tartarated Antimony, as recommended in former editions of the *Companion*

Dose.—10 to 30 minims = 0.6 to 1.8 cc, as an emetic, 2 to 4 fl drms = 7.1 to 14.2 cc

Contains 1 grain in 240 minims

Foreign Pharmacopœias—Official in Belg, Dutch, Ger and Jap (Vinum Stibiolum), 1 in 250, Mex (Vino estibiado), 1 in 300, Span. (Vino Emetico), 1 in 250, Russ (Vinum Stibio-Kalii Tartarici), 1 in 250, US (Vinum Antimonii), 1 in 250, all with Sherry Austr (Vinum Stibii Kalio-Tartarici), 1 in 250, Hung (Vinum Stibiato-Tartaricum), 1 in 240, Swiss (Vinum Stibiolum), 1 in 250, all with Malaga Wine Port (Vinho Antimonial), 1 in 200 of Port Wine All by weight, except US Not in Dan, Fr, Ital, Norw or Swed

Tests.—Antimonial Wine has a specific gravity of about 1.006; it contains about 4.75 p c w/v of total solids and about 19 p c w/v of Absolute Alcohol

Not Official

UNGUENTUM ANTIMONII TARTARATI—Tartarated Antimony, in fine powder, 1, Simple Ointment, 4—*B P* 1885
This has been incorporated in the *B P C*

ANTIPYRINE.

See PHENAZONUM.

Not Official.

APIOL.

An oily liquid, with a peculiar odour and disagreeable taste, obtained from the Fruits of *Apium Petroselinum*, L (Parsley)

Medicinal Properties—It is useful in amenorrhœa and dysmenorrhœa

Dose.—3 to 5 minims = 0.18 to 0.3 cc

Prescribing Note—Usually given in capsules

Foreign Pharmacopœias—Official in Fr, Mex and Port (Apiol), Dan. and Norw (Ætheroleum Petroselinii), Swed includes Fructus Petroselinii, and an Aqua of it.

Apiol was described by Messrs Joret and Homolle, who introduced the substance into medicine, as a yellow, oily, non volatile liquid, but the Apiol obtained by us from the Homolle capsules, although yellow in colour, was volatile in the vapour of Water to the extent of 95 p c. Witney went into the subject in 1880, and describes Apiol as an impure Essential Oil of Parsley containing minute quantities of soft resin, and the Apiol of Homolle as the Essential Oil of Parsley Seeds with small traces of a soft resin. The Essential Oil of Parsley is a yellow, oily liquid, and as such has been made official in the Danish and Norwegian Pharmacopœias.

Arising out of a discussion as to what should be the colour of liquid Apiol, it was suggested in *C D* '94, 11 17, that it was simply an alcoholic extract of Parsley Seeds, but this product is green, and contains but a small proportion (under 15 p c) of the Essential Oil of Parsley.

The stearoptene from the Oil is known as **Crystallised Apiol**

Not Official.

APOCYNUM

Syn—CANADIAN HEMP

The dried Rhizome of *Apocynum Cannabinum*, L., is official in U S

Medicinal Properties—It has been used in the United States for some years as a cardiac tonic, and diuretic in cardiac dropsy. Also as a **fluid extract** (dose 5 to 15 minims = 0.3 to 0.88 c c) in pleurisy with effusion.

It also possesses emetic and cathartic properties, but as it is a drastic purgative it should be given with some caution.

The diuretic action of Canadian Hemp was favourably considered, although it was admitted that it might produce violent emesis and catharsis. These undesirable results were, however, attributed to the admixture of the bitter fibre of the wood with the bark of the root—*B M J* '97, 11 1714.

In 1 minim doses has been successfully employed (*L* '05, 11 955) in a case of incompetency and ascites, increased by 2 minims up to 10 minim doses 3 times daily, and subsequently reduced to 6 minims.

The most irritant of all cardiac tonics, and not a drug to be employed advantageously in medicine—*B M J* '06, 11 1460.

Descriptive Notes—The root is usually $\frac{1}{8}$ to $\frac{1}{2}$ inch (4 to 6 mm) thick, cylindrical, somewhat angular, and longitudinally wrinkled with a few transverse fissures. It has an orange-brown bark, which becomes grey brown on keeping, whitish or pinkish internally, nearly as thick as the woody portion, and contains large laticiferous vessels. The yellowish wood has several concentric rings, is finely radiate and coarsely porous, taste bitterish and somewhat acrid, it has scarcely any odour. The root of *A. androsaemifolium*, L., is sometimes confused with it, but it has a white porous wood and groups of stone cells in the outer part of the bark.

FLUIDEXTRACTUM APOCYNII—100 of Apocynum in No. 60 powder is moistened with 40 of a mixture of Glycerin 10, Alcohol (95 p c) 60, and Water 30, packed in a percolator, then enough menstruum added to saturate the powder and leave a stratum above it, macerated for 48 hours, and percolation allowed to proceed slowly, gradually adding, first, the remainder of the menstruum, afterwards, a mixture of Alcohol (95 p c) 60 and Water 40, until the Apocynum is exhausted, reserving the first 90 of percolate and evaporating the remainder at a temperature not exceeding 50° C (122° F) to a soft extract which is dissolved in the reserved portion, and enough menstruum added to make 100. Average Dose—1 c c (15 minims)—*U S P*.

This has been incorporated in the *B P C*.

TINCTURA APOCYNII—Root, 1, Alcohol (60 p c), 10, by maceration.

Dose,—5 to 10 minims = 0.3 to 0.8 c c

This has been incorporated in the *B P C*.

APOCYNIN—An amorphous resinous substance, almost insoluble in Water, but soluble in Alcohol (90 p c) and in Ether.

APOMORPHINÆ HYDROCHLORIDUM.

APOMORPHINE HYDROCHLORIDE

FR, CHLORHYDRATE D'APOMORPHINE, G&S, APOMORPHINHYDROCHLORID;
ITAL, CHLORIDRATO DI APOMORFINA, SPAN, CLORURO DL APOMORFINA

$C_{17}H_{17}NO_2, HCl$, eq 301.36

White or greyish-white, odourless, minute, shining, needle-shaped crystals, which should be kept from the light in dark amber-coloured, thoroughly dry, glass bottles, and protected as far as possible from the air.

Apomorphine is obtained from Morphine by the abstraction of a molecule of Water, and Apomorphine Hydrochloride is prepared by its Hydrochloride in sealed tubes with an excess of strong Zinc Chloride. The *B P* mentions Morphine or Codeine, but the product of the abstraction of a molecule of Water from Codeine is generally considered to be Apocodeine.

Solubility—About 1 in 60 of Water, 1 in 50 of Alcohol (90 p c); nearly insoluble in Chloroform and in Ether, 1 in 100 of Glycerin.

The solubility in Water is given in *B P* as 1 in 50, minimum quantity of Water required for complete solution in 3 days at 60° F is between 1 in 56 and 1 in 60, but if dissolved by the aid of a gentle heat it will remain in solution at 1 in 50. The aqueous solution, on being gently warmed, rapidly turns green.

The material used was recrystallised, air-dried, and powdered. It lost 3 p c of hygroscopic moisture on heating in a water-bath, which was exactly regained after 12 hours' exposure to air.

The solubility of Apomorphine Hydrochloride has been stated (*P J* '05, i, 230, *C D* '05, i, 282) to be 1 in 53 of Water, and 1 in 48 of Alcohol (90 p c). The *Companion* figures have been since shown to be quite correct in references to them in *P J* '06, i, 345, *C D* '06, i, 471.

Medicinal Properties.—The most reliable emetic, $\frac{1}{10}$ grain hypodermically, or $\frac{1}{2}$ grain by the mouth, usually acts promptly (2 or 3 minutes) without the production of much preceding nausea or depression, or unpleasant after-effects. As a hypodermic injection in cases of poisoning, especially if unable to swallow, and if emesis be indicated.

Invaluable as an expectorant in acute and chronic bronchitis with viscid secretion, and in croup, in bronchial irritation due to inhalation of factory dust, and in asthma.

As a hypnotic, $\frac{3}{8}$ grain hypodermically. As patients are occasionally susceptible to emetic action it is well to begin with $\frac{2}{10}$ grain, as a hypnotic—*L*, '00, i, 1088.

Absolutely inert as an emetic in alcoholic poisoning—*L*, '00, i, 1685.

In the acute stage of alcoholism, delirium tremens, 5 drops of the Injection are generally sufficient to produce, within 5 minutes, several hours' sleep. This dose can be repeated if necessary, the patient ought to be in horizontal position, occasionally, vomiting precedes the sleep. It is not a remedy for alcoholic craving—*B M J* '07, ii, 951.

As a hypnotic, hypodermically, in acute alcoholism, it has not received the recognition it deserves—*M R* '07, ii, 144.

Dose.— $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.003 to 0.007 gramme, by hypodermic injection, by the mouth, $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.007 to 0.016 gramme.

Ph Ger maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme.

Prescribing Notes.—*Its aqueous solution on keeping, or on being gently warmed, rapidly turns green. This green coloration is said to be due to the liberation of free Apomorphine by the alkalinity of the glass, and can be prevented by adding a few drops of dilute Hydrochloric Acid to the preparation. The official injection keeps fairly well for a month or so. Some authorities are of opinion that the Ammonia in the an causes the alkalinity.*

Official Preparation—Injectio Apomorphinæ Hypodermica

Not Official.—Hypodermic Discs, Haustus Apomorphinæ Compositus, Mistura Apomorphinæ et Terebenti, Pastillus Apomorphinæ et Codeinæ, and Syrupus Apomorphinæ Hydrochloridi

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U S

Tests—The distinguishing tests of Apomorphine Hydrochloride are the production of a white precipitate, rapidly changing to green on exposure to air, when its aqueous solution is made alkaline with Potassium or Sodium Bicarbonate. The changed alkaloid is soluble in Alcohol (90 p c) producing an emerald green solution, in Chloroform producing a fine violet tint, and in Ether a purple coloration. An aqueous solution affords a precipitate with Mercuric-potassium Iodide (Mayer's) Solution, with Iodo-potassium Iodide (Wagner's) Solution, and with Mercuric Chloride Solution. The two following tests serve to distinguish Apomorphine Hydrochloride from Morphine on the one hand, and from Codeine, Nacaine, and Narcotine on the other. The first is given in the *B P*, but without any explanation of its object, the second is peculiar to the *U S P*.

(1) Ferric Chloride Test—solution colours Apomorphine Hydrochloride Solution red, whilst Morphine yields a dull greenish-blue coloration.
(2) A solution of 0.05 gramme of Apomorphine Hydrochloride when shaken with a solution of 0.05 gramme of Ferrous Sulphate in 10 c c. of Water, gradually acquires a blue colour, changing after some time to a bluish-black, the original blue should be restored upon the addition of Alcohol (90 p c), but Carbon Bisulphide, Chloroform and Ether should remain colourless when shaken with the aqueous liquid.

An aqueous solution gives with Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, on the addition of Ammonia Solution, the precipitate dissolves, but its solution is immediately reduced.

A number of colour reactions for the identification of the alkaloid are given in small type below, and where possible comparison is made with statements appearing in the respective Pharmacopœias.

Apomorphine Hydrochloride and Pilocarpine Nitrate are two exceptions to the usual *B P* requirements, that the alkaloidal salt shall afford the tests distinctive of the individual acid radical entering into its composition.

The more generally occurring impurities are acidity, the presence of salt which has undergone decomposition, and mineral matter. Acidity is covered by the requirements of the three Pharmacopœias that the salt or its aqueous solution shall be neutral, or only very feebly acid to Litmus paper or solution. A salt which has been kept under unfavourable conditions, or which has been in stock for a long while, almost invariably contains a considerable amount of salt which

has undergone change. The *BP*, *USP* and *PG* state that a salt, one part of which with 100 parts of Water yields an emerald green solution should be rejected, the *PG* further stating that a colourless or at most only a pale reddish liquid should result on shaking the dry salt with Ether. Neither the *BP* nor *USP* mention the likelihood of mineral matter being present, but *PG* states that the salt shall leave no residue on ignition.

Colour Tests—With Nitric Acid it yields a blood red coloration *BP*, and *PG*, fading to orange *USP*, with dilute Solution of Ferric Chloride it gives a deep red coloration *BP* and *USP* (distinction from Morphine which yields a blue colour).

USP also includes the following colour reactions which do not appear in *BP* or *Ger*.

Sulphuric Acid does not colour the salt, but with Sulphuric Acid containing (a) a trace of Selenious Acid it produces a dark blue colour fading to violet and then to pink, (b) a trace of Ferric Chloride, a pale blue colour, (c) a trace of Ammonium Vanadate, a violet blue colour, changing to deep greenish-blue, (d) a little Paraldehyde, a green colour, fading to reddish-brown, (e) Potassium Iodate, a black colour, changing to brown and finally to pale brown, (f) a trace of Nitric Acid, a blood red colour, fading to orange. A crystal of Apomorphine Hydrochloride and a crystal of Potassium Nitrate with Sulphuric Acid added are coloured red, and on stirring with a glass rod the solution becomes green, then blue, then purple, and finally cherry red. Acetic Acid dissolves the salt without colour, but on adding a trace of Potassium Iodate, the solution turns blood red, changes to purple, and on adding a little Ether and shaking, the latter assumes a blue colour. Gold Chloride T.S. produces a reddish purple precipitate in a solution of the salt.

Preparation.

INJECTIO APOMORPHINÆ HYPODERMICA. HYPODERMIC INJECTION OF APOMORPHINE

Apomorphine Hydrochloride, 1 gram, Diluted Hydrochloric Acid, 1 minim, Distilled Water (recently boiled), *q. s.* to make 110 minims.

5 minims = $\frac{1}{2}$ grain Apomorphine Hydrochloride

Dose.—5 to 10 minims = 0.3 to 0.6 c.c.

Not Official.

DISCS OF APOMORPHINE— $\frac{1}{10}$ to $\frac{1}{20}$ grain dissolved in 6 to 10 minims of Distilled Water at the time or using subcutaneously—*St. Bartholomew's*.

PASTILLUS APOMORPHINÆ ET CODEINÆ—Apomorphine Hydrochloride, $\frac{1}{2}$ grain, Codeine, $\frac{1}{10}$ grain—*Martindale and B.P.C.*

HAUSTUS APOMORPHINÆ COMPOSITUS—Apomorphine Hydrochloride, $\frac{1}{10}$ grain, Diluted Hydrochloric Acid, 1 minim, Oil of Turpentine, 10 minims, Mucilage of Gum Acacia, *q. s.*; Spirit of Ether 10 minims, Distilled Water, to 1 fl. oz.—*Muddeler*.

MISTURA APOMORPHINÆ ET TEREBINI—Apomorphine Hydrochloride, $\frac{1}{10}$ grain, Pure Terebene, 15 minims, Peru Balsam, 10 minims; Mucilage Mixture, to 1 oz.—*Guy's*.

SYRUPUS APOMORPHINÆ HYDROCHLORIDI—Apomorphine Hydrochloride, 5 grains, Diluted Hydrochloric Acid, 10 minims, Alcohol (90 p.c.), 7 fl. drm., Distilled Water, 7 fl. drm., Syrup, to produce 20 fl. oz. Dissolve the salt in the Spirit and Water mixed, then add the Acid and the Syrup—*B.P.C. Formulary 1901*.

Dose— $\frac{1}{2}$ to 1 fl. drm. = 1.8 to 3.6 c.c.

Contains $\frac{1}{32}$ grain in 1 fl drm

This has been incorporated in the *B.P.C.* as follows —

Syrupus Apomorphinæ—Apomorphine Hydrochloride, 0·05, Diluted Hydrochloric Acid, 0·25, Alcohol (90 p c), 4·50, Distilled Water, 4·50, Syrup, q s to make 100

This contains about $\frac{1}{32}$ grain in 1 fl drm

AQUÆ.

WATERS

The Waters of the British Pharmacopœia, all of which are distilled, except Aqua Camphoræ and Aqua Chloroformi, are as follows, the formulas are given under the names of the substances from which they are prepared —

AQUA ANETHI From the dried ripe fruit
 AQUA ANISI From dried ripe Anise fruit
 AQUA AURANTII FLORIS From the flowers Imported
 AQUA CAMPHORÆ
 AQUA CARUI From the dried fruit
 AQUA CHLOROFORMI
 AQUA CINNAMOMI From the bark
 AQUA DESTILLATA
 AQUA FENICULI From the dried ripe fruit
 AQUA LAUROCERASI From fresh leaves
 AQUA MENTHÆ PIPERITÆ With oil and distilled
 AQUA MENTHÆ VIRIDIS With oil and distilled
 AQUA PIMENTÆ From the dried unripe fruits
 AQUA ROSÆ From the fresh flowers
 AQUA SAMBUCI From the fresh flowers Imported

In preparing distilled aqueous liquids only good, natural, potable Water must be employed, as directed for 'Distilled Water'

In India and other tropical countries the Waters of Anethum, Anisum, Carum, Cinnamomum, Fœniculum, Mentha Piperita, Mentha Viridis, and Pimenta, may be prepared from the oils without distillation, using 1 of Oil and 2 of Calcium Phosphate to 500 of Distilled Water

AQUA DESTILLATA.

DISTILLED WATER

FR, EAU DESTILLÉE, GER, DESTILLIRTWASSER, ITAL, ACQUA DISTILLATA, SPAN, AGUA DESTILADA

A clear, colourless, odourless, tasteless, neutral, limpid liquid, obtained by distilling good natural Water of a potable quality

Foreign Pharmacopœias.—Official in all

Tests—Distilled Water at the normal temperature and pressure boils at 100° C (212° F), and should evaporate leaving scarcely a weighable residue. It should possess neither taste nor odour, and should be perfectly neutral. It may contain as impurities various metals, dissolved solids, Chlorides, Nitrates, Nitrites, Sulphates, organic matter, and Ammonia, all of which are specially mentioned in the *B.P.* The *USP* and *PG* examine for heavy metals. The *PG* and *USP* include a test for Carbonic Acid gas. The quantities used

by the *B.P.* (25 cc) and by the *P.G.* (10 cc) for the determination of the amount of solid residue are ridiculously small, the *USP* quantity (1000 cc) is much more to the point, and admits of a quantitative determination. The tests for Chlorides, Nitrates, Nitrites, and Sulphates given in the *B.P.* are those mentioned in the Appendix, but with the exception of those for Chlorides and Sulphates are too crude unless specially applied. The *USP* employs the Diphenylamine test for Nitrates, requiring that no blue colour shall be produced, and the Sulphanilic Acid and Naphthylamine Acetate test for Nitrites, stipulating that no pink coloration shall be produced in 5 minutes. The *B.P.* test for Ammonia is carried to the other extreme, and is described with more minuteness than its importance justifies. The *B.P.*, *USP* and *P.G.* employ 100 cc of Distilled Water in carrying out the test for readily oxidisable organic impurities with Potassium Permanganate Solution, but the relative quantities of Sulphuric Acid and Potassium Permanganate vary, as does also the time of boiling. The *B.P.* and *P.G.* tests coincide pretty closely, with the exception of the relative strengths of the diluted Sulphuric Acid official in each Pharmacopœia. Both boil for 3 minutes, and at the end of this time the liquid should retain its pink colour. The *USP* boils for 10 minutes, and requires in addition that the pink colour should not wholly disappear if the vessel is covered to protect it from dust and set aside in a dark place for 10 hours.

A detailed comparison of the tests adopted by the three Pharmacopœias for the detection of the above impurities is for convenience made in small type below.

Residue—The *B.P.* requires that the residue left on the evaporation of 25 cc should be scarcely visible, the *P.G.* states that 10 cc of Distilled Water when evaporated should not leave a weighable residue, and the *USP* that 1000 cc of Distilled Water, after evaporation to dryness on a water bath, should not leave more than 0.050 grammes of residue.

Silver Nitrate—It should not be affected by Silver Nitrate T.S., indicating the absence of Chlorides, *P.G.* and *USP*.

Mercuric Chloride—It should not be affected by solution of Mercuric Chloride, *P.G.*

Barium Chloride—It should not yield the slightest turbidity on the addition of Barium Chloride 1:5, indicating the absence of Sulphates, *USP*.

Ammonium Oxalate—No turbidity should result on the addition of Ammonium Oxalate T.S., indicating the absence of Calcium, *USP*.

Hydrogen Sulphide—It should not be affected by Hydrogen Sulphide T.S., even after the addition of Solution of Ammonia, *P.G.* It should not respond to the time-limit test for heavy metals, *USP*.

Calcium Hydroxide—When mixed with twice its volume of Lime Water it should remain clear, indicating the absence of Carbonic Acid, *P.G.* and *USP*.

Diphenylamine—If 10 cc of Distilled Water mixed with a few drops of Diphenylamine T.S. be carefully poured upon about 3 cc of Sulphuric Acid free from Nitrous compounds, contained in a test-tube, so as to form a separate layer, no blue colour should be formed at the line of contact of the two liquids, *USP*.

Sulphanilic Acid and Naphthylamine Acetate—If to 50 cc of Water contained in a glass cylinder 2 cc each of Sulphanilic Acid T.S. and Naphthylamine Acetate T.S. are added, and the solution well mixed, no distinct pink coloration should appear within 5 minutes, if the cylinder be placed upon a white surface and viewed from above, *USP*.

Mercuric Potassium Iodide—The *BP* requires that it shall not contain more than 0.005 part per million, equivalent to 0.00035 grain per gallon, of Ammonia as determined by the colour produced when 2 c.c. of Potassium mercuric Iodide (Nessler's) Solution is added to 100 c.c. of the Water, the colour is compared with that produced by 0.25 c.c. of Nessler's Ammonium Chloride Solution diluted with 50,000 c.c. of Ammonia free Water. A somewhat similar test is given in the *USP*, which uses 50 c.c. of Distilled Water and 2 c.c. of reagent. This should not yield a yellow or brownish tint when contained in a glass cylinder placed on a white surface and viewed from above. The *USP* does not, however, direct the use of a comparative test.

Potassium Permanganate—If 100 c.c. of Water be heated to boiling with 1 c.c. of Diluted Sulphuric Acid (the *USP* uses 10 c.c. of Diluted Sulphuric Acid) and solution of Potassium Permanganate be added and the mixture boiled for 3 minutes (10 minutes *USP*), the liquid should not be decolorised, *BP*, *USP* and *PG*. The *BP* states that the liquid should retain its colour for 1 hour, and the *USP* that the colour should not be completely destroyed by boiling for 10 minutes, nor should it wholly disappear if the vessel be afterwards set aside in a dark place and covered for 10 hours. The quantity of Potassium Permanganate Solution ordered in the *BP* is 0.1 c.c. of a mixture of 1 part solution of Potassium Permanganate (1 p.c.), and 2 parts of Water, in the *PG* 0.8 c.c. of a 0.1 p.c. w/w solution of the salt is used, and in the *USP* 0.1 c.c. of a Tenth normal Volumetric Solution is used.

ARAROA.

ARAROA

BP Syn—GOA POWDER, CRUDE CHRYSAROBIN

Fine powder, or in more or less agglomerated particles, yellow when first obtained, but rapidly becoming of a dull ochre or brown colour. Obtained from fissures in the trunk of *Andra Araroba*, Aguiar and freed from woody fragments.

Descriptive Notes—The crude drug of commerce is usually of an umber-brown colour but yellowish internally and more or less mixed with, or attached to, pieces of the heart wood of the tree. From these pieces it should be freed and then dried and powdered, and it may then vary in colour from brownish-yellow to umber-brown. Samples which have been imported in a damp condition and exposed to atmospheric Ammonia become deep brown or purple. It varies much in the amount of wood it contains. Good specimens yield as much as 80 p.c. of soluble matter to Benzol or Chloroform.

Official Preparation.—Used to prepare Chrysarobin.

Tests.—Araroba should be soluble in hot Chloroform to the extent of not less than 50 p.c., and when the solution is filtered and evaporated to dryness, the powdered residue is known as Chrysarobin.

Not Official

ARECA.

The Seed of the *Areca Catechu*, L., the betel nut tree. Imported from the East Indies.

Medicinal Properties—Astringent, narcotic, anthelmintic. A remedy for tapeworm. 60 grains = 4 grammes, of powdered Areca Nut made into a ball.

with Honey, answers well as a vermifuge for a large dog. A paste is made of the powder for a dentifrice

Areca Nut Charcoal used also as a dentifrice

Foreign Pharmacopœias—Official in Ger and Swiss, Semen Arecae

Descriptive Notes—The seeds of the Areca palm vary a good deal in size and somewhat in shape. In size they vary from 1 to $1\frac{1}{2}$ inches (25 to 37 mm) in diameter in the broadest part, the transverse section is white marbled with brown, owing to the infolding of the seed coat in the albumen. The taste is astringent and slightly acid. The hemispherical or rounded fruits, derived from the variety *alba*, are much less active as a vermifuge than the conical or typical form. The larger and more mature seeds are to be preferred to the smaller for medicinal use.

ARECOLINÆ HYDROBROMIDUM—Fine white needles, readily soluble in Water and in Alcohol (90 p c), difficultly soluble in Ether and in Chloroform

Sialagogue, diaphoretic and anthelmintic. $\frac{1}{2}$ p c solution applied to the eye produces tingling followed by myosis. Maximum effect in from 10 to 15 minutes, lasts about one hour. Administered internally causes vomiting and diarrhoea—*B M J* '99, 1 82

Dose— $\frac{1}{16}$ to $\frac{1}{8}$ grain = 0.00625 to 0.00125 gramme, to be given with caution.

Foreign Pharmacopœias—Official in Fr, Ger, Swed and Swiss.

Tests—It should have a melting point of 167°C (332.6°F); *Ph Codex* gives 170°C (335°F). A 1 in 10 aqueous solution of the salt yields a brown precipitate with Iodine Solution, a brown precipitate with Bromine Water, and a light yellow precipitate with Silver Nitrate Solution, but none with Platinum Chloride Solution, Mercuric Chloride Solution, or Tannic Acid Solution. It should leave no residue when ignited with free access of air.

ARGENTI NITRAS.

SILVER NITRATE

B P Syn—LUNAR CAUSTIC

AgNO_3 , eq 168.69

Fr, AZOTATE D'ARGENT, GER, SILBERNITRAT, ITAL, NITRATO DI ARGENTIO, SPAN, NITRATO ARGENTICO CRISTALIZADO

Colourless, transparent, tabular, rhombic crystals. It should be kept in well stoppered, dark amber-coloured vials protected from the light and dust.

Solubility.—100 grains in 50 minims of Water, measuring 80 minims, 1 in 18 of Alcohol (90 p c). Insoluble in strong Nitric Acid.

Medicinal Properties.—Astringent, sedative, antispasmodic. It is useful in hæmatemesis, gastric ulcer, diarrhoea and cholera, as well as in chronic nervous irritability of and pain in the stomach; also in some nervous diseases, as epilepsy, chorea and locomotor ataxy. It is employed in chronic constipation as an **enema**, 60 grains dissolved in 60 oz of Water, after clearing away the contents of lower bowel, and as a **bougie** in chronic gonorrhoea. A dark line on the edges of the gums, removable by a course of Acid Tartrate of Potassium, precedes the indelible discoloration of the skin and mucous membranes (argyria), produced by the long-

continued internal administration of this salt Its administration should be interrupted for fourteen days at the end of two or three months, however small the dose More than 100 grains per month should not be given

Externally as a local stimulant to weak and callous ulcers, fistulæ, and aphthous affections of the mouth, as a caustic to warts and poisoned wounds As a local application to prevent pitting in smallpox, and to relieve the itching in pruritus, it is also applied, under Cocaine, to ulcers of the cornea 1 to 3 grains to the oz is employed for lotions and collyria, in all forms of conjunctivitis and both as a prophylactic and curative in ophthalmia neonatorum, and as an injection in urethritis, cystitis, and vaginitis For eczema or pityriasis of the ear, a 1 in 20 solution in Spirit of Nitrous Ether answers well Hæmostatic for leech bites

Chilblains are sometimes painted with a strong solution of Silver Nitrate

A weak solution (1 in 500) for obstinate forms of eczema in children —*L M R* '88, 525

In cholelithiasis —*B M J E* '02, 1 99

Powdered Silver Nitrate in antral empyema —*B M J E* '90, 1 96

A 2 p c solution of Silver Nitrate by far the best prophylactic in ophthalmia neonatorum —*B M J* '03, 11 135, *L* '03, 11 163, a 4 grains to the oz solution for treating the same, *Pr lxxv* 561, as a prophylactic, 1 p c is even more efficacious than the 2 p c advocated by Ciede, whilst the inflammatory reaction is less marked, *L* '07, 11 588

Intravenous injection of a 2 or 5 p c solution in the treatment of septic conditions —*B M J E* '02, 1 12

Strong Solution of Potassium Iodide, or Potassium Cyanide, has been suggested for the removal of the black stains on the skin produced by Silver Nitrate

Dose — $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.016 to 0.032 gramme

Prescribing Notes — *Prescribed in pills with Massa Kaolin. Solutions should be dispensed in stoppered bottles*

For application to the skin, a solution in Spirit of Nitrous Ether has been recommended This solution throws down a light coloured precipitate, but does not itself become black like a simple spirituous solution It, however, blackens the skin in a shorter time.

Incompatibles — The Alkalis and their Carbonates and Alkaloids, all Bromides, Chlorides, Iodides, and Phosphates, Solutions of Arsenic, and Tannin

Official Preparations — Argenti Nitras Induratus and Argenti Nitras Mitigatus Used in the preparation of Argenti Oxidum

Not Official — Mild Caustic Points, Argenti Iodidum Nascens, Argentum Foliatum, Actol, Albaigin, Argentamin, Argentol, Argonin, Argyrol, Collargol, Ichthaigau, Itrol, Largin, Novaigan, Protargol, and Tachiol

Antidotes — Aqueous solution of Common Salt, Milk or some demulcent drink given freely, Emetic, White of Egg

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Span, Swed, Swiss and U S Also fused Nitrate of Silver in all except Belg, Dan, Fr, Ger, Ital, Swed and Swiss Russ now includes only a fused Nitrate.

Tests. — The distinguishing tests for Silver Nitrate are that its aqueous solution yields with Hydrochloric Acid or a soluble Chloride a white curdy precipitate insoluble in Nitric Acid but soluble in Ammonia Solution, Potassium Chromate Solution yields a red precipitate disappearing on the addition of a solution of a soluble Chloride,

Bromide or Iodide, and destroyed by mineral acids, a solution of Ferrous Sulphate gently poured on to a well-cooled mixture of its aqueous solution and Sulphuric Acid, produces a brown or purplish-brown coloration at the junction of the two fluids, an aqueous solution warmed with Sulphuric Acid and a strip of Copper liberates red fumes. It is officially required to yield 84.3 p.c. of Silver Chloride as ascertained gravimetrically by precipitating 1 gramme of the salt with Hydrochloric Acid, the filtrate from the precipitate, when evaporated to dryness should leave no residue. The *P.G.* completely precipitates the Silver from 5 cc of an aqueous 1 to 20 solution by the addition of an excess of Hydrochloric Acid, and requires that the evaporated filtrate should leave no residue; the *USP* uses a 1 to 10 solution and requires that not more than 0.1 p.c. of residue should remain. The *BP* gravimetric test indicates 100.0 p.c. of the pure salt, the *USP* volumetric test indicates 99.98 p.c. of the pure salt, neither a gravimetric nor a volumetric determination is given in *P.G.* The *USP* test is performed with Deci-normal Volumetric Sodium Chloride Solution, the excess being titrated back with Deci-normal Volumetric Silver Nitrate Solution.

The more generally occurring impurities are Copper, Iron and Lead, Potassium, Sodium and Sulphates, all of which are mentioned in the *BP*. Copper and Lead are the more important, the former is readily detected by the blue colour of the ammoniacal solution, the latter by the insoluble Sulphate thrown down on the addition of Sulphuric Acid.

Sulphuric Acid—5 cc of an aqueous solution (1-10) with 20 cc of hot diluted Sulphuric Acid and heated to boiling should give no white precipitate (absence of Lead), *USP*.

Volumetric Determination—0.5 gramme of Silver Nitrate is dissolved in 10 cc of Distilled Water and well mixed with 30 cc Tenth-normal Volumetric Solution of Sodium Chloride and 3 drops of Potassium Chromate T.S. when not more than 0.4 cc Tenth-normal Volumetric Silver Nitrate Solution should be required to impart to the liquid a permanent red colour, *USP*.

Preparations.

ARGENTI NITRAS INDURATUS. TOUGHENED CAUSTIC.

Silver Nitrate, 19, Potassium Nitrate, 1. Mix by fusion.

Foreign Pharmacopœias—Belg (*Argentum Nitricum Stylis*), Fr. (*Crayons d'Azotate d'Argent*), Ital (*Nitrato d'Argento Fuso con Nitrato di Potassio*), Mex (*Lápices de Nitrato de Plata*), Span (*Nitrato Argentico Mitigado*), all Silver Nitrate 9, Potassium Nitrate 1.

Argentum Nitras Fusus.—4 grammes of Hydrochloric Acid are added to 100 grammes of Silver Nitrate in a porcelain dish and melted at as low a temperature as possible, stirred well and poured into suitable moulds, which should be kept in dark amber-coloured vials protected from light—*USP*.

ARGENTI NITRAS MITIGATUS. MITIGATED CAUSTIC.

Silver Nitrate, 1, Potassium Nitrate, 2. Mix by fusion.

Foreign Pharmacopœias—Official in Austr, Ger., Russ and Swiss, 1 in 3, Jap, 1 in 2, all (*Argentum Nitricum c. Kalio Nitrico*). Dan (*Nitras Argenticus bis Mitigatus*), 1 in 3, Norw and Swed (*Nitras Argenticus Mitigatus*), 1 in 3, Fr. (*Crayons d'Azotate d'Argent*

Mitigé), containing $\frac{1}{2}$, $\frac{1}{3}$, and $\frac{1}{4}$ of Nitrate of Silver, US (*Argentum Nitras Mitigatus*), 1 in 8 Not in the others

Mild Caustic Points, made by fusing Potassium Nitrate in various proportions with Silver Nitrate, are used by oculists and others

Not Official

ARGENTUM FOLIATUM (*Austr, Belg, Dan, Dutch, Fr, Ger, Russ and Swiss*)—Thin leaves of pure Silver, which dissolve in Nitric Acid, yielding a clear colourless solution

ARGENTI IODIDUM NASCENS—Freshly precipitated Silver Iodide has been recommended in conjunctival catarrhs

Silver Iodide is a heavy, amorphous, yellowish powder, which should be kept in dark amber-coloured vials, protected from light

ACTOL (Silver Lactate)—A white amorphous powder, or as colourless crystalline needles Soluble 1 in 20 of Water Introduced as an antiseptic Useful as an injection ($\frac{1}{4}$ to $\frac{1}{2}$ grain per oz) in gonorrhoea The injection is attended with some pain

Possesses no advantage for ophthalmic work over Silver Nitrate—*B M J* '01, 11 1383

ALBARGIN (Gelatose Silver, Silver Glutin)—Bright yellow powder, soluble in Water Stated to be an active antiseptic—*P J* '01, 11 345

ARGENTAMIN—Silver Phosphate dissolved in Ethylenediamine solution Antiseptic and astringent A dilution of 1 to 4000 of Water has been recommended for urethral injection in gonorrhoea As a 5 p c solution in ophthalmic work—*B M J E* '95, 11 20, '96, 11 64, *L* '95, 11 47, *B M J* '01, 11 1333

ARGENTOL—A compound of Silver with Oxymolinol A sparingly soluble yellowish powder, recommended as an antiseptic application to wounds and ulcers—*P J* '97, 1 369, '98, 11 342

ARGONIN—Is obtained by precipitating Silver Nitrate and Casein-soda with Alcohol Contains about 4 p c of Silver It is a fine white powder, soluble in Water It is recommended as a disinfectant—*P J* (3) xxv 1193, *J S C I* '95, 1060, *L* '95, 11 47 A 2 p c aqueous solution gradually increased to 10 p c, recommended in the treatment of gonorrhoea—*B M J E* '96, 11 64, *T G* '97, 740, *B M J* '01, 11 1333

ARGONIN L—Contains 10 p c Silver and is readily soluble 1 p c solution used successfully in the treatment of anterior and posterior urethritis—*B M J E* '99, 1 96

ARGYROL—Fine black glistening hygroscopic scales readily reduced to a powder Readily soluble in Water forming a dark brownish black solution, but insoluble in Alcohol (90 p c) It contains 30 p c of Silver combined with a protein extracted from wheat 5 p c injections in acute gonorrhoea—*L* '03, 11 1716 Instillations of 5 to 50 p c solutions are useful in ophthalmic work, and are painless

In the treatment of ophthalmia neonatorum 1 or 2 drops of a 20 to 50 p c solution put into each eye is stated (*P* lxxv 561) to be certainly less irritating than, and quite as effective as, Silver Nitrate Bacteriological experiments have shown that Argyrol is more effective in killing micro organisms than Silver Nitrate in the strengths in which the latter can be tolerated in the eye

In the form of 'drops' of a 25 p c solution it is of the greatest value in checking suppuration from the conjunctiva, and in some most unpromising cases it proved to be the best remedy—*L* '05, 1 1416

An ointment containing 10 grains to the oz of Vasoline is simply invaluable (*M P* '05, 11 138) in eczematous conjunctivitis or keratitis complicated with marked photophobia, blepharospasm and watering of the eyes, and when the photophobia is very intense 2 to 4 grains of alkaloidal Atropine and a similar amount of alkaloidal Cocaine may with advantage be added to the prescription

Of the three Silver compounds, Protargol, Collargol and Argyrol, the latter

(*L* '06, ii 14) gives the best results in acute conjunctivitis. It is the least irritating and possesses quite a wonderful power in lessening discharge and relieving pain. It can be used in as great a strength as 50 p.c., and may be applied as an ointment or solution. It is so non-irritating that it can safely be injected into the anterior chamber to control intraocular suppuration. It is the most valuable remedy we possess in the treatment of purulent ophthalmia, either in the adult or in the new-born child. It is also useful in blenorrhagia of the lachrymal passages.

Uses and limitations in eyework of Argylol.—Great advantage, even in strong solution, over Silver Nitrate, lies in the fact that its action is painless, and that it can be used freely by the patient without risk.—*Am. J. Ophth. and Surg. Jour.* '07, i 396. In ophthalmia neonatorum, solutions of Argylol are much better than strong solutions of Silver Nitrate, because they are practically inert, and therefore do not injure the conjunctival epithelium (*B. M. J.* '07, ii 67a), as an enema, in ulcerative colitis (*L* '04, ii 1209), yields good results applied as 20 p.c. solution in ophthalmia neonatorum (*B. M. J.* '04, ii 1246), 25 p.c. solution applied in acute conjunctival cases, and great improvement followed application in a case of persistent ophthalmia.—*B. M. J.* '04, ii 1633.

See also Summary given below

COLLARGOL (Colloid Silver)—Black or greyish-black shining scales, with a metallic lustre. Soluble 1 in 2 of Water. Antiseptic and disinfectant. Employed in the form of a 15 p.c. ointment. Intravenously as an injection, 5 to 20 c.c. of a $\frac{1}{2}$ to 1 p.c. solution. As a 1 to 5 solution in ophthalmic work.—*L* '02, ii 1800, *B. M. J.* '01, ii 95, '02, ii 16, *M. P.* '02, i 85, *P. J.* '02, i 11.

Of the three Silver compounds, Protargol, *Colloidal Silver*, and Collargol is to be preferred (*L* '06, ii 14) in recent wounds of the eyeball, and is employed for the most part in the form of Gelatin waters containing 10 p.c. of the drug. It is also used in the form of solution and as an ointment, the strength varying from 5 to 20 p.c.

ICHTHARGAN (Silver Thiohydrocarbauo sulphonate, Silver Ichthyolate)—A light brown, odourless, amorphous powder, containing 30 p.c. Silver. Soluble 1 in 6 of Water and in Glycerin, Insoluble in Alcohol (90 p.c.) and in Ether. Powerful antiseptic. Useful as an injection, 0.02 to 0.2 p.c., in gonorrhoea. 1 to 3 p.c. solution in affections of the posterior urethra.—*B. M. J.* '01, ii 104, '02, ii 16, '03, ii 31, *P. J.* '01, ii 299.

ITROL (Silver Citrate)—A white odourless powder, containing about 63 p.c. of Silver. Only slightly soluble in Water (1 in 4000). Antiseptic. Useful in gonorrhoea. As an injection (1 in 8000 to 1 in 4000 solution). As an insufflation. As a dusting powder for wounds. In the form of sticks for fistulae, deep wounds, and endometritis.—*Itrol* 2 to 5, White Wax, 10. Ol. Theobromi, 9, melt and divide into 80.—*P. J.* '96, i 243, '97, ii 254, *Pr. ix* 292, *B. M. J.* '00, i 99, *T. G.* '99, 631, *P. J.* '99, ii 185, *B. M. J.* '01, ii 1333.

Official in Swed

LARGIN (Silver Albuminate)—A light brown, amorphous odourless powder. Soluble 1 in 8 of Water. Contains 11 p.c. Silver. Introduced as an antiseptic. Useful in gonorrhoea as an injection (1 in 4000). It is stated to have given very satisfactory results in superficial eye-diseases, such as acute infectious ophthalmia in 8 to 10 p.c. solution. Inferior to the Nitrate or Protargol in gonorrhoeal ophthalmia. Even saturated solution stated to cause no pain.—*B. M. J.* '00, i 622, *B. M. J.* '00, i 68, *P. J.* '00, i 413.

NOVARGAN—A fine yellow odourless powder, yielding neutral red-brown solutions with Water, which should be kept in non-actinic glass bottles. An antiseptic in gonorrhoea.

PROTARGOL (Silver Protein).—A light brown or yellow, odourless powder, possessing a disagreeable metallic taste. It should be preserved in well stoppered glass bottles of a dark amber tint and protected as far as possible from the light. It is soluble 1 in 2 of Water. A powerful antiseptic and germicide possessing deep penetrating powers, and stated not to precipitate albumen. Has been recommended in $\frac{1}{2}$ to 2 p.c. solution as an unirritating and successful injection in gonorrhoea.—*B. M. J.* '97, ii 98, '98, i 40, '98, ii 2, *Pr. ix* 292.

and 311, *L* '97, ii, 1628, '98, i 872, *B M J E* '99, ii 56, 59, *L* '02, i 1525, *B M J E* '99, i 24, '01, ii 103, *B M J* '07, i 912, *FT* '07, 46 Irritant effect sometimes seen probably due to worthless imitations of Protargol, or to the solutions not being made up freshly and with cold Water—*B M J E* '07, i 72 In conjunctival affections in the form of a 5 to 10 p.c. solution—*B M J E* '99, i 8, *T G* '99, 318, *Pr* lxxiv 476, *L* '99, ii 1046, '01, ii 553 In gonorrhoeal ophthalmia, 10, 20, and 80 p.c. solutions may be used—*L* '01, ii 553, *B M J* '01, ii 1333 1 p.c. solution never fails to cure—*B M J* '07, i 912 Zinc Sulphate stated to be incompatible with Protargol—*P J* '02, ii 293 In phthisis, injection of 40 c.c. of saline solution containing $1\frac{1}{2}$ to 2 $\frac{1}{2}$ grains Protargol, preceded and followed by an injection (through the same needle) of a few c.c. of pure saline solution—*Proc Brit Cong on Tuberc* iii 444, *L* '01, ii 309 $1\frac{1}{2}$ p.c. solution as an injection, and local application of a 10 to 20 p.c. solution to diseased area in urethral hæmorrhage in gonorrhoea—*L* '02, i 1526 Two cases of argyrosis following the use of Protargol in conjunctivitis—*L* '02, ii 1199 Injections of solutions of this salt, followed by astringents—*e g*, Zinc Chloride or Acetate—are considered (*L* '04, ii 1223) best in the treatment of gonorrhoea 2 to 5 p.c. solutions are usually employed—*Pr* lxxiv 225

Of the three Silver salts, Protargol, Collargol and Argylol, in chronic inflammation of the conjunctiva and the edges of the eyelids, Protargol, in from 10 to 25 p.c. solution or ointment brushed vigorously over the affected parts, produces far quicker and better results than are to be obtained by any other known method of treatment—*L* '06, ii 13

See also Summary below

Prescribing Notes—In preparing solutions of Protargol it is preferable to place the greater portion of the Water which is intended to be used into a vessel and then to introduce the Protargol, which dissolves gradually, finally adding the rest of the Water. Shaking should be avoided until solution is complete

Foreign Pharmacopœias—It is official in Austr., Belg., Jap. and Swiss

Tests—A 1 p.c. aqueous solution of Protargol is not precipitated by the addition of alkalis, Sodium Chloride Solution, or by Ammonium Hydrosulphide Solution. Hydrochloric Acid produces a precipitate soluble in excess of the reagent, and Picric Acid Solution yields a yellow precipitate. Solutions of Protargol yield the Biuret reaction when mixed with an equal volume of Potassium or Sodium Hydroxide Solution and a drop or two of a diluted Copper Sulphate Solution. It should leave on ignition about 8 p.c. of metallic Silver which should conform to the characteristic tests and freedom from impurities mentioned under 'Silver Nitrate'

Liquor Protargol—Protargol, 40, 80, 120, or 160 grains, Distilled Water, to 1 fl. oz.—*London Ophthalmic*

TACHIOI (Silver Fluoride)—Colourless, transparent crystals, changing rapidly on contact with air. Readily soluble in Water, and the solutions, if not too strong, are permanent. Introduced as a new antiseptic. It is stated to possess powerful bactericidal powers, superior to Carbolic Acid, and only slightly inferior to Corrosive Sublimate. Employed in from a 1 in 1000 to 1 in 5000 solution to disinfect cavities and suppurating sinuses, in tubercular lesions, and ulcerative processes. It is also stated to have met with some success in ophthalmic practice, its non-irritating character being an advantage. Its solution, however, blackens linen fabrics—*L* '02, i 993, ii 1707, *C D* '02, i 309 A 1 in 8000 to 1 in 5000 solution has been shown (*B M J E* '05, i 24) to be an excellent antiseptic and antifermentative for the stomach, whether used for washing out or given internally

Summary of Experiments with Organic Silver Compounds.—A series of experiments were carried out to determine the bactericidal action of Silver compounds, and the results were embodied in a report to the British Medical Association—*B M J* '06, ii 359 The various Silver compounds investigated fall into three groups (1) Those which are powerfully bactericidal, (2) one (Nargol) much less powerfully bactericidal, (3) two (Argylol and Collargol) which possess practically no bactericidal action whatever. The first group includes most of the substances investigated, namely, Silver Nitrate, Silver Fluoride, Actol,

Itrol, Argentamin, Argonin, Ichthargan, Largin, Novargan and Protargol. These in solutions containing the same percentage of combined Silver is closely similar, and it is practically impossible to place them in any order of activity which would be true under all circumstances.

As Argyrol and Collargol are not bactericidal, it is evident that the amount of Silver which a compound may contain is no criterion of its bactericidal power. Moreover, in view of the results obtained with Argyrol, it seems impossible to attribute the good effects which many clinicians have obtained with it to its bactericidal action.

F. 1475, largely confirmatory of the above—*B M J* '07, ii

Silver Nitrate in eye diseases is a dangerous remedy, except in the hands of the surgeon, and its application even in a 2 p.c. solution is attended with such excruciating pain as to make the strongest shudder—*Scot Med and Surg Jour.* '07, i 396.

ARGENTI OXIDUM.

SILVER OXIDE.

Ag_2O , eq 230.10.

FR, OXYDE D'ARGENT, GÉR, SILBEROXID

A brownish-grey odourless powder when freshly prepared, but becoming of a blackish-brown colour on drying or on exposure to the air.

When mixed with readily oxidisable substances great heat is evolved, and the mixture is liable to explode.

It may be prepared by the interaction of solutions of Silver Nitrate and Calcium Hydroxide.

It should be preserved in dark amber-tinted dry bottles, and protected from dust and ammoniacal fumes.

Medicinal Properties—It has the general therapeutic qualities of the Nitrate, without its escharotic effect. It is more slowly absorbed, and is said to be less liable to discolour the skin.

Dose.— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 grammes.

Prescribing Notes—Usually given in a pill, made with *Massa Kaolini*.

It prescribed with Creosote or with the Chlorides in pills, the Oxide must be first diffused through some inert matter such as Kaolin, as the heat produced in rapidly reducing the Silver or by its combining with it causes the mass to become red-hot, or to explode.

Incompatibles—Bromides, Chlorides, and Iodides. Organic and readily oxidisable matter.

Foreign Pharmacopœias—Official in U.S. Not in the others.

Tests.—The distinguishing tests for Silver Oxide are the ease with which it undergoes decomposition when mixed with readily oxidisable substances such as Creosote and Potassium Permanganate, such decomposition being accompanied with rapid disengagement of heat, and if in a confined space with explosive violence, the evolution of Oxygen when heated to dull redness leaving a residue of metallic Silver, which should, when dissolved in Nitric Acid, yield the tests distinctive of Silver given under that substance. It is

officially required to indicate 100.0 p.c. of pure Silver Oxide as gravimetrically determined by solution in Nitric Acid and precipitation with Hydrochloric Acid, 1.237 grammes of Silver Chloride should be yielded by 1 gramme of Silver Oxide. The *USP* gravimetric test indicates 99.8 p.c. of pure Silver Oxide, which is equivalent to 92.9 p.c. of pure metallic Silver, determined as the residue left on igniting the Oxide in a porcelain crucible, 0.5 gramme should not leave less than 0.464 gramme. Silver Oxide is not official in the *PG*.

The more generally occurring impurities are metallic Silver, Lead, Copper, and Iron, all of which are mentioned in *BP*. The presence of metallic Silver is evidenced by the evolution of reddish fumes when the Oxide is dissolved in Nitric Acid. Lead and Copper are detected as described under *Argentum Nitratum*. The *USP* includes a test for Carbonate and a limit of Chloride, requiring that the Oxide should dissolve without effervescence in Nitric Acid, and that when a weighed quantity of 0.2 of a gramme of the Oxide is dissolved in 1 c.c. of Nitric Acid mixed with twice its volume of Water, and 10 c.c. of Ammonia Solution are added, the liquid diluted to 60 c.c. the addition of 1 c.c. of Nitric Acid to 10 c.c. of this dilution should not produce a cloudiness.

ARMORACIÆ RADIX.

HORSERADISH ROOT

FR, RAIFORT, GER, METTRICH, ITAL, RAPANO RUSTICANO,
SPAN, RABONO RUSICANO

The fresh Root from cultivated plants of *Cochlearia Armoracia*, L. Most active in the autumn and early spring before the leaves have appeared.

Medicinal Properties—Sialagogue, stomachic, slightly diuretic, and diaphoretic. Used in atonic dyspepsia and as a condiment, also as a sudorific in chronic rheumatism. Externally as a rubefacient. The infusion is used as a gargle.

Official Preparation—*Spiritus Armoracæ Compositus*

Not Official—*Infusum Armoracæ Compositum*, *Sirup de Raifort Composé*

Foreign Pharmacopœias—Official in Dutch, Fr, Port (*Rabao Rustico*), Span and Mex. Not in the others.

Descriptive Notes—The root is imported from Belgium in barrels, that usually sold not being cultivated in this country. It is made up in bundles of about twenty roots. It is whitish or yellowish-white externally, about 8 to 12 inches (20 to 30 cm.) or more long, about an inch (2.5 cm.) in diameter, cylindrical, slightly enlarged and marked with leaf scars at the crown, odourless, until scraped or broken, when it gives off a pungent odour, the taste is very pungent. Aconite root has on rare occasions been mistaken for it, but the latter is conical, tapering much below, blackish-brown externally and not pungent. Horseradish root is usually sold in the fresh state, rarely in a dried condition.

Preparation.**SPIRITUS ARMORACIÆ COMPOSITUS.** COMPOUND SPIRIT OF HORSE RADISH.

Scraped Horseradish Root, 1 oz, Dried Bitter-Orange Peel, 1 oz, Nutmeg, 11 grains, Alcohol (90 p c), 5 fl oz, Distilled Water, 6 fl oz Mix and distil 8 fl oz

Dose.—1 to 2 fl dm = 3 6 to 7 1 c c

Foreign Pharmacopœias—Not in the other Pharmacopœias Port, compound Wine, Mex (Alcoholato de coquealia); Span (Alcohol de coquealia) They all differ widely from the above

Not Official.

INFUSUM ARMORACIÆ COMPOSITUM—Fresh Root, sliced, 1; Black Mustard Seed, 1, Compound Spirit of Horseradish, 1, boiling Distilled Water, 20 Macerate two hours, strain, and add the Spirit

It is found in practice that a temperature of 150° to 180° F makes the strongest infusion

Dose.—1 to 2 fl oz = 28 4 to 56 8 c c, as a warm stimulant.

SIROP DE RAIFORT COMPOSÉ (Antiscorbutic Syrup)—Cochlearia Leaves, 1000, Water-cress Leaves, 1000, Horseradish Root, 1000, Dried Leaves of Menyanthes Trifoliata, 100, Bitter-Orange Peel, 200, Ceylon Cinnamon, 50, White Wine, 4000, White Sugar, 5000, all by weight Bruise the Water-cress and Cochlearia Leaves, cut up the Horseradish Root, the Leaves of Menyanthes Trifoliata and the Bitter-Orange Peel, break up the Cinnamon Bark Macerate the whole in the White Wine for two days and distil on a water-bath Collect 1000 of the aromatic liquor, and with it prepare a syrup in a closed vessel on the water-bath in the proportion of 100 grammes of Sugar for each 100 grammes of the liquor—*Fr*

ARNICÆ RHIZOMA.**ARNICA RHIZOME.***B P Syn*—**ARNICÆ RADIX.**

FR., RACINE D'ARNIQUE, **GER.**, ARNIKAWURZEL, **ITAL.**, RHIZOMA DI ARNICA; **SPAN.**, RIZOMA DE ARNICA.

The dried Rhizome and Roots of *Arnica montana*, Linn.

Collected in the mountainous parts of Central and Southern Europe

The dried flower-heads are official in the *Ind.* and *Col Add*

Medicinal Properties.—Stomachic and slightly stimulant, irritant to the stomach and bowels in large doses The **tincture** is used externally for bruises and sprains, diluted with Water, but inflammation of the skin may be set up, equally good results have been produced by the application of Spirit and Water.

Official Preparation—*Tinctura Arnicæ*

Not Official—*Arnica Opodeldoc*, *Extractum Arnicæ Radicis Fluidum*, *Linimentum Arnicæ*

Antidotes—Opium, Morphine

Symptoms of poisoning by Arnica are violent vomiting, intense headache, diarrhoea, colic, feeble pulse

Foreign Pharmacopœias—Official in Austr, Ital and Port, root and flowers, Hung, root, leaves, and flowers, Belg, Dan, Dutch, Fr, Ger, Jap, Norw, Russ, Swed, Swiss and U S, flowers, Mex, rhizome, leaves, and flowers, Span, rhizome and flowers

Descriptive Notes—The rhizome as imported requires to be picked over before use, and freed from various foreign matters and vegetable fibres, which are almost always mixed with it. It is officially described as about 1 to 2 inches (2½ to 5 cm) in length, ½ to 1 inch (4 to 6 mm) in diameter, cylindrical and curved, horizontal. It is often furnished with one or more terminal proliferations, giving it a jointed appearance, and with rather distant, wavy, unbranched, brittle roots, on the lower surface only, and is sometimes terminated at the crown by the hairy bases of the leaves. The taste is acid and bitter, and the odour characteristic, recalling that of apples. The transverse fracture shows a greyish or dirty white centre, and resin ducts near the inner margin of the cortex. It is sometimes mixed with the rhizome of species of *Hieracium*, or of *Geranium urbanum*, Linn. The former lacks the resin ducts and odour, and the latter has a clove odour, and astringent taste, and has roots all round the erect rhizome or rootstock.

In the *PG* and *USP* the flowers are official instead of the root, but these are liable to attacks by insects, and lose their volatile oil more quickly than the rhizome.

Preparation

TINCTURA ARNICÆ TINCTURE OF ARNICA

1 of Arnica Rhizome in No 40 powder, percolated with Alcohol (70 p c) to yield 20 (1 in 20)

Foreign Pharmacopœias—Official in Fr and U S, 1 in 5, Belg, Dan, Dutch, Ger, Norw, Russ, Swed and Swiss, 1 and 10, Port 1 in 10, all from flowers, Port, 1 in 5, from the root, Ital, root 1, Alcohol (60 p c) 10, Austr, root 4, flowers 1, Alcohol (70 p c) 25, Hung, root 6, leaves 3, and flowers 1, dilute Alcohol (70 p c) 50, Mex, dried leaves 1 in 5, Span, root 1, flowers 1, Alcohol (70 p c) to 10, all are by weight except U S.

Tests—Tincture of Arnica has a specific gravity of 0.890 to 0.895, contains about 0.6 p c w/v of total solids and about 68 p c w/v of Absolute Alcohol.

TINCTURA ARNICÆ FLORUM—Percolate 1 of Arnica flowers with Alcohol (45 p c) to make 10.

Official in the *Ind* and *Col Add* for North American Colonies

Not Official

ARNICA OPODELDOC—White Soap, 4, Alcohol (90 p c), 10, Tincture of Arnica, 5, Camphor, 1. Dissolve by heat, and strain.

This has been incorporated in the *BPC* under the title **Linimentum Arnicæ**.

EXTRACTUM ARNICÆ RADICIS FLUIDUM—1 in 1, made percolation with a mixture of Alcohol 8, Water 1.—*USP* 1890.

This has been incorporated in the *BPC*.

Not Official

ARSENII BROMIDI LIQUOR.LIQUOR POTASSII ARSENIATIS ET BROMIDI (*U S N F*) CLEMENS' SOLUTION

115. Potassium Bicarbonate, 587 grains, Bromine, 235 grains, Water, *q s*. Boil the Arsenic and Potassium Bicarbonate in 8 oz of Water till dissolved, when cold add 16 oz of Water, then the Bromine, and make up with Water to 32 fl oz. Stir occasionally during a few hours, then filter.

This Liquor was originally described by Dr. Clemens as 'a chemical union of Arsenic and Bromine,' but as the action of Bromine on Arsenious Acid results in the formation of Arsenic Acid and Hydrobromic Acid, the above formula has been adjusted to yield these products as Potassium salts.

The solution contains Arsenic equal to 1 p.c. of Arsenious Anhydride Recommended in the treatment of diabetes.—*J. M. R.* '83, 86

Dose —1 to 5 minims = 0.06 to 0.3 c.c.

ARSENII IODIDUM.

ARSENIOS IODIDE

 AsI_3 , eq 452.20*Syn* —ARSENIC IODIDE

Orange or orange red coloured crystals, having a faint odour of Iodine, and which lose Iodine on exposure to air and light.

It should be kept in dark amber-coloured well stoppered glass bottles in a cool place.

Solubility —1 in 11 of Water, 1 in 42 of Alcohol (90 p.c.); 1 in 19 of Carbon Bisulphide.

It is gradually decomposed by boiling Water and by boiling Alcohol.

Medicinal Properties.—Has been used in obstinate cutaneous affections of syphilitic and tubercular origin.

Dose — $\frac{1}{8}$ to $\frac{1}{2}$ grain = 0.0034 to 0.013 gramme.

Prescribing Notes —It is generally given as Donovan's Solution, or in a pill well triturated with Milk Sugar and massed with Glucose.

Official Preparation —Liquor Arsenii et Hydragryri Iodidi.

Foreign Pharmacopœias —Official in Jap, Mex (*Yoduro de Arsenico*) and U.S. Not in the others.

Tests —The distinguishing tests for Arsenious Iodide are that its aqueous solution shall, when acidulated with Hydrochloric Acid, yield on addition of Hydrogen Sulphide Solution a yellow precipitate soluble in Potassium or Sodium Hydroxide Solution, in Potassium Carbonate Solution or in Ammonium Carbonate Solution, being reprecipitated on the addition of Hydrochloric Acid, a cold porcelain vessel impinged upon the flame of the ignited gas produced by the interaction of Zinc, Hydrochloric Acid and a little of an aqueous solution of the salt, acquires a dark metallic looking stain readily.

dissolved by Chlorinated Soda Solution, Hydrogen Arsenide is evolved when a little of the solution is boiled with Zinc and Potassium or Sodium Hydroxide Solution, this gas yields a black stain to filter paper moistened with Silver Nitrate Solution, and a dark metallic stain on porcelain as above when ignited, Stannous Chloride Solution containing a large excess of Hydrochloric Acid yields a brown or brownish black coloration when mixed with a little of an aqueous solution, an aqueous solution acidulated with Hydrochloric Acid yields a dark metallic looking deposit on Copper when boiled with that metal, the Copper when removed, dried between folds of bibulous paper, and heated in a clean, dry test-tube yields a sublimate of white characteristic octahedral crystals. The aqueous solution yields with Silver Nitrate Solution a curdy yellow precipitate insoluble in Nitric Acid, almost insoluble in Ammonia Solution, but soluble in Potassium Cyanide solution, with a small quantity of Chlorine Solution and Starch Mucilage it yields an intense blue coloration, disappearing on heating and reappearing as the liquid cools, when treated with Chlorine Solution and shaken with Carbon Bisulphide, the latter liquid assumes a fine violet coloration. No mention of a volumetric or gravimetric method of determination is made in the *BP*, but the *USP* requires it to contain not less than 16.3 per cent of Arsenic as indicated by titration with Deci-normal Volumetric Iodine Solution by the method described in the small type below.

The more generally occurring impurities are free acid, either Arsenous or Hydriodic, and mineral matter. The neutrality of the aqueous solution to Litmus affords an indication of the former, the latter is revealed by a residue remaining after volatilisation.

Volatilisation—When heated on a water bath no loss in Iodine occurs, *USP*, but at a higher temperature it volatilises, *BP* and *USP*. Violet vapours of Iodine being set free, *BP*.

Volumetric Determination—0.5 gramme of Arsenious Iodide and 2 grammes of Sodium Bicarbonate dissolved in 50 c.c. of Water require not less than 21.9 c.c. Tenth normal V.S. of Iodine to impart a slight yellow tint to the solution, *USP*.

Preparation

LIQUOR ARSENII ET HYDRARGYRI IODIDI. SOLUTION OF ARSENIOS AND MERCURIC IODIDES

Arsenious Iodide, 87½ grains, Mercuric Iodide, 87½ grains, Distilled Water, *qs* to make 20 fl. oz. (1 of each in 100)

A yellowish, odourless liquid, with a disagreeable metallic taste, sp. gr. 1.015 to 1.018. It is also known as **Donovan's Solution**.

Dose.—5 to 20 minims = 0.3 to 1.2 c.c.

11 minims contain $\frac{1}{10}$ grain of each salt.

Incompatibles—Acids, the salts of Morphine, or other Alkaloid, and Corrosive Sublimate.

Foreign Pharmacopœias—Official in U.S., 1 in 100,

ASAFETIDA.

ASAFETIDA

FR, ASAFETIDA, GER, ASANT, ITAL, ASSAFETIDA, SPAN, ASAFETIDA

A gum resin obtained by incision from the Root of *Ferula foetida*, Regel, and probably other species

Medicinal Properties—Nervine stimulant, expectorant, laxative and calmative Useful in cases of flatulence, in hysterical paroxysms, also in some forms of chronic bronchitis, very useful as an enema in the flatulent distension of typhoid or peritonitis, and in infantile convulsions

As a successful preventive against abortion—*M A* '93, 64, *B M J B* '95, 1 35

Dose.—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—In pill massed with a little dilute Alcohol They are best varnished, as silver leaf is affected by this drug The Tincture may be prescribed with Aromatic Spirit of Ammonia, or with the Tinctures of Valerian and Hyoscyamus When diluted with water to form a mixture, it requires the addition of Mucilage of Gum Acacia

Official Preparations—Tinctura Asafetide Used in the preparation of Pilula Aloes et Asafetidae, Pilula Galbani Composita, and Spiritus Ammoniae Fetidis

Not Official—Enema Asafetida, Mistura Asafetida Composita, Pilula Asafetida

Foreign Pharmacopoeias—in Austr, Belg, Dan., Dutch, Fr, Ger., Hung., I, J, N, S, S, Russ, Span, Swed, Swiss and U. S.

Descriptive Notes—The Asafetida of commerce varies exceedingly in appearance and in purity It may occur in whitish tears from $\frac{1}{2}$ inch (12.5 mm) in diameter, or in flattened tears up to 3 inches (75 mm) or more in diameter, or in masses of agglomerated tears of various sizes, either whitish or brownish or reddish-brown according to age, and more or less agglutinated with darker gum resin and mixed with stones More rarely balls of sand, about 4 cm. in diameter, made with the liquid gum resin are offered under the name of stony Asafetida As it is very difficult to purify without loss of essential oil, it is very necessary that the purest drug obtainable should be selected for dispensing purposes. It is officially limited to the rounded or flattened tears, more or less agglutinated, which are dull yellow in colour and darken on keeping, and internally yellowish and translucent, or milky-white and opaque, the freshly exposed surfaces gradually assuming a pink colour, changing to red and finally to reddish-brown The *U S P.* requirements are much the same The *P G* gives *Ferula Nardus*, Boiss, also as a source of the drug The gum resin of that species, however, never becomes red on keeping, and Arclison ascertained that in Kashmir it is not collected. For the purpose of being powdered the *P G* directs that it should be dried over quicklime, and triturated at the lowest practicable temperature The Asafetida in tears that enters into commerce does not invariably turn red on exposure Even fine-looking specimens in the form of tears, sometimes contain half their weight of pebbles

covered with a thin layer of Asafetida, and it is only from selected specimens, easily distinguished by their light weight, that a drug affording only 10 p c of ash can be obtained. Usually the tears met with in commerce yield only one-third of their weight of this quality. Indeed, in this as in other cases, where much impurity is present, a purified preparation extracted by Alcohol from the crude drug should be official for dispensing purposes.

Tests—The distinguishing tests for Asafetida are that when triturated with Water it forms a milky-white emulsion, which assumes a yellowish coloration on the addition of a few drops of Ammonia Solution, the addition of Nitric Acid diluted with an equal volume of Water to the freshly fractured surface produces a greenish coloration, when strongly heated in a dry test-tube, cooled and treated with boiling Water it yields a liquid which, when largely diluted and made alkaline with Ammonia Solution, exhibits a blue fluorescence. This test is known as the umbelliferone test, and the remarks on the test appearing under Ammoniacum apply equally here. It is officially required to contain not less than 65 p c of matter soluble in Alcohol (90 p c), but only the best quality will yield this proportion. The *USP* and *PG* fix 50 p c as the limit of matter soluble in Alcohol (90 p c). The method adopted by the *PG* of weighing the insoluble residue is preferable to determining the amount dissolved, on account of the loss of volatile constituents during evaporation. The determination of the Acid, Ester and Saponification values affords useful indications of the purity, but no mention of them appears in the *BP*. Commercial Asafetida has an Acid value of from 60 to 80, an Ester value of from 80 to 130, and a Saponification value of 120 to 185. A sample of fine selected tears examined in the author's laboratory, which yielded 3.1 p c of ash and contained 68.3 p c of matters soluble in Alcohol (90 p c), had an Acid value of 131.9, an Ester value of 119.3, and a Saponification value of 251.3. Another sample which contained 37.8 p c of ash, and 44.5 p c of matters soluble in Alcohol (90 p c), had an Acid value of 43.4, an Ester value of 127.4, and a Saponification value of 170.8.

The more generally occurring impurities are inferior varieties of Gum and mineral matter. The presence of the former is detected by the Alcohol-solubility, the latter by the ash. A lengthy controversy has taken place over the amount of ash permissible. An ash limit of 20 p c has been suggested (*YBP* '00, 405), and it has been stated (*CD* '99, 11 983) that at that date it was practically impossible to obtain any considerable quantity which would satisfy the *BP* tests. On the other hand, it was stated (*CD* '99, 11 1037) that provided a fair price were paid a sufficient supply of *BP* quality could readily be procured. The *USP* and *PG* both state not more than 10 p c. The *Dutch Pharmacopœia*, which in the Third Edition adopted a 20 p c limit, in the new Fourth Edition allows not more than 10 p c.

Preparations

PILULA ALOES ET ASAFETIDÆ, 1 in 4. *See* ALOES.

PILULA GALBANI COMPOSITA. about 1 in 3½ See GALBANUM

SPIRITUS AMMONIÆ FETIDUS, about 33 grains in 1 oz.
See AMMONIA.

TINCTURA ASAFETIDÆ. TINCTURE OF ASAFETIDA.

1 of Asafetida, maceiated with Alcohol (70 p c), to yield 5
(1 in 5)

Dose — ½ to 1 fl drim = 18 to 36 c c

Foreign Pharmacopœias Official in Belg (1 and 4½), Dan, Dutch, Fr, Ital, Jap, Norw, Port, Span, Mex, Swed and Swiss, 1 and 5, U S, 1 in 5, all by weight, except U S, not in Austr, Ger, Hung or Russ

Tests — Tincture of Asafetida has a specific gravity of about 0.912. It contains about 9½ p c w/v of total solids and about 60 p c w/v of Absolute Alcohol

Not Official

ENEMA ASAFÆTIDA.—Asafetida, 30 grains, Distilled Water, 4 fl oz. Rub the Asafetida in a mortar with the Water added gradually so as to form an emulsion — *B P* 1885

Tincture of Asafetida, 1 fl drim, Mucilage of Starch, 1 fl oz — *St. Thomas's*.

This has been incorporated in the *B P C* as follows —

Tincture of Asafetida, 3, Mucilage of Starch, q s to make 100.

MISTURA ASAFETIDA COMPOSITA — Asafetida, picked, 5 grains, Liquid Extract of Cascara Sagrada, 10 minims, Ammonium Carbonate, 1 grain, Infusion of Valerian (1 in 40), to 1 fl oz. The Ammonia in this mixture develops the taste and odour of the other constituents — *St. Thomas's*.

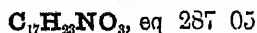
This has been incorporated in the *B P C*, using 5 grains of Ammonium Carbonate in place of 4

PILULÆ ASAFÆTIDA — Asafetida, 20 grammes, Soap, in fine powder, 6 grammes, Water, q s to make 100 pills — *U S P*. Each pill contains 3 grains of Asafetida

This has been incorporated in the *B P C*

ATROPINA.

ATROPINE



FR, ATROPINE, GER, ATRORIN, ITAL, ATROPINA, SPAN., ATROPINA.

Colourless, odourless acicular crystals, or as a white more or less amorphous powder. Taste bitter and acrid. Obtained from the leaves and root of Belladonna. It gradually assumes a yellowish tint on exposure to air, and should be kept in well-stoppered dark amber-tinted bottles

It is officially regarded as an alkaloid, obtained from Belladonna Leaves or Root. The bulk of the alkaloid existing in Belladonna is Hyoscyamine, which is isomeric with Atropine, and the former has a constant tendency to change into the latter

Atropine (uncombined with an Acid) easily decomposes when heated. A solution of 1 in 200 of Water heated in a basin or on a water-bath for two hours was so completely decomposed that it lost

its alkaline reaction and ceased to precipitate with Mercuric Chloride Solution, after eight hours the reaction was faintly acid

Solubility —1 in 500 of Water, 1 in 3 of Alcohol (90 p c), 1 in 25 of Ether, 1 in 1 of Chloroform, 1 in 52 of Glycerin, 1 in 15 of Oleic Acid

Medicinal Properties —The Ointment is used for the relief of pain arising from muscular spasm, and for neuralgia. See also Atropinæ Sulphas and Belladonna

Dose — $\frac{1}{200}$ to $\frac{1}{100}$ grain = 0.0003 to 0.0006 gramme

It is freely soluble in Oleic Acid, and is sometimes applied as a 1 or 2 p c solution

Official Preparation —Unguentum Atropinæ

Not Official —Atropinæ Oleas, Unguentum Atropinæ, Unguentum Atropinæ Dilutum, Unguentum Atropinæ cum Acido Borico, Unguentum Atropinæ cum Cocainæ, and Atropinæ Oleas

Antidotes —In case of poisoning by Atropine, the antidotes are the same as for Belladonna

Foreign Pharmacopœias —Official in Fr, Mex, Port, Span and U S Not in the others

Tests —The distinguishing tests for Atropine are the melting point which, when pure, should be about 115.5°C (248°F), the Aurichloride, which should be dull and pulverulent in physical appearance, should possess a melting point of 137°C (278.6°F), the strongly marked mydriatic action when a drop of a very dilute solution is carefully instilled into the eye, the optical inactivity, the purple-violet coloration produced when a crystal is evaporated to dryness with fuming Nitric Acid, and the residue is moistened with a freshly prepared alcoholic Potassium Hydroxide solution, this test being known as Vitali's test, the yellow precipitate changing to red, which occurs when an alcoholic solution is warmed with Mercuric Chloride Solution, the alkaline reaction to Litmus and Phenolphthalein, which distinguishes Atropine and its isomers from almost all other known alkaloids. The *BP* mentions most of these tests, but not the optical inactivity or melting point of the Aurichloride, although it describes the method of producing the salt and its physical appearance and includes a melting point for Hyoscyne Aurichloride. The *USP* states that pure Atropine Aurichloride melts at 136°C (276.8°F). The colour reaction with Vitali's test is very delicate, and although other alkaloids, Pseudoaconitine, Veratrine, and Strychnine, afford colours somewhat resembling this, there is no difficulty in detecting Atropine in the pure state when unmixed with other alkaloids. The *USP* states that the presence of Strychnine masks this reaction. The latter Pharmacopœia also states that it produces no precipitate with Platonic Chloride Solution and that it yields a pink coloration not dissipated by 0.5 gramme of Chloral Hydrate, when a crystal is treated with Sulphuric Acid containing a drop of Cresol, which distinguishes it from most other alkaloids. It is also stated to give a peculiar odour indicative of a mixture of Rose, Orange flower, and Melilot

when heated with a little Sulphuric Acid, the odour changing to one of Bitter Almonds on the addition of a small crystal or two of Potassium Bichromate

Atropine may be readily titrated with Deci- or Centi-normal Volumetric Hydrochloric or Sulphuric Acid Solution, using Cochineal Solution as an indicator. 1 cc Deci-normal solution = 0.0287 gramme of Atropine, and 1 cc of Centi-normal solution = 0.00287 gramme of Atropine. Atropine is not official in the *P.C.*

The more generally occurring impurities are mydriatic alkaloids other than Atropine, *e.g.*, Hyoscyamine and Scopolamine, Morphine, and mineral matter

Hyoscyamine and Scopolamine are readily detected by their optical activity, Morphine, if present, by the red coloration produced when a small quantity of the alkaloid is treated with a mixture of Sulphuric and Nitric Acids, mineral matter by the amount of residue left on ignition, which should be nil

Gold Chloride—The Chloraurate may be made by adding Gold Chloride T.S. to a solution of Atropine in dilute Hydrochloric Acid, washing, collecting, and drying the precipitate, which should be yellow and lustreless, *U.S.P.*

Vitali's Test—If Atropine be moistened with Nitric Acid (fuming, *B.P.*), and heated in a porcelain dish on a water-bath to dryness, the residue, yellow, (*U.S.P.*) gives with alcoholic Potassium Hydroxide solution, reddish-violet coloration, *B.P.*, an intense violet coloration, *U.S.P.* (which is due to a fragment of the base). The base directs the addition of a fragment of the base to the solution. Hyoscyamine and Hyoscyamine give a reddish-violet coloration, the presence of Strychnine will mask the coloration, *U.S.P.*

Sulphuric Acid—Sulphuric Acid when added to Atropine should show no colour (absence of readily carbonisable organic impurities), nor should subsequent addition of Nitric Acid yield any colour (absence of and discolouration from Morphine), *U.S.P.*

Preparation

UNGUENTUM ATROPINÆ.—ATROPINE OINTMENT.

Atropine, 2; Oleic Acid (by weight), 8; Lard, 90. (1 in 50)

Not Official.

UNGUENTUM ATROPINÆ—Atropine, 4 grains, Soft Paraffin, 1 oz.; heat till dissolved and stir till cold—*London Ophthalmic*; *Middlesex*, same strength with Vaseline

UNGUENTUM ATROPINÆ DILUTUM—Atropine 1 p.c. in fine powder, incorporated in Yellow Soft Paraffin of uniform consistency having a melting point of about 35° C—*St. Thomas's*

This has been incorporated in the *B.P.C.*

UNGUENTUM ATROPINÆ CUM ACIDO BORICO.—Atropine, 4 grains, Powdered Boric Acid, 60 grains, Soft Paraffin, 1 oz—*London Ophthalmic*

UNGUENTUM ATROPINÆ CUM COCAINA—Atropine, 4 grains, Cocaine, 8 grains, Soft Paraffin, 1 oz, heat till the alkaloids are dissolved—*London Ophthalmic*

Atropine, 1 p.c., Cocaine, 2 p.c.—*St. Thomas's*

This has been incorporated in the *B.P.C.*

ATROPINÆ OLEAS—Atropine, 8 grains; Oleic Acid, 1 oz.

OLEATUM ATROPINÆ.—Atropine, 2, Alcohol (95 p c), 2, Oleic Acid, by weight, 50, Olive Oil, *q s* to make 100 by weight. Triturate the Atropine in a tared mortar with the Alcohol, then add an equal volume of the Oleic Acid and, after warming the mortar, stir until the Alcohol has evaporated, add the remainder of the Oleic Acid and continue stirring until the Atropine is dissolved, then add Olive Oil to make 100 by weight —*U S P*

This has been incorporated in the *B P C* with the title **Oleinatum Atropinæ** *Syn.* OLEATUM ATROPINÆ

HOMATROPINE —*See* p 598

ATROPINÆ SULPHAS

ATROPINE SULPHATE

FR, SULFATE D'ATROPINE, GER, ATROPINSULFAT, ITAL, SOLFATO DI APROINA,
SPAN, SULFATO DL ATROPINA



A white, or almost white, odourless, more or less crystalline powder, having a bitter, nauseous taste

It is the Sulphate of an alkaloid obtained from *Belladonna Leaves* or *Root* and may be obtained by neutralising Atropine with Diluted Sulphuric Acid. As obtained commercially it almost invariably contains a small proportion of Hyoscyamine Sulphate

Solubility—10 in 4 of Water, 1 in 4 of Alcohol (90 p c) Insoluble in Ether and Chloroform

Medicinal Properties—Mydriatic, anhidrotic, antigalactagogue. Employed locally to dilate the pupil and paralyse the accommodation, in iritis, and before testing refraction or making ophthalmoscopic examination, used also to cause retraction of protruding iris, as it increases intraocular tension it does harm in glaucoma. It is frequently combined with Morphine in hypodermic administration to prevent the undesirable effects of the latter. Injected as near the nerve as possible in sciatica, hypodermically in ovarian and uterine pain. The hypodermic method is also the best to diminish the sweating of phthisis, for which purpose, in doses of $\frac{1}{100}$ to $\frac{1}{1000}$ grain, Atropine is very useful, it at the same time relieves the cough, or 1 or 2 minims of the *Liquor Atropinæ Sulphatis* may be given by the mouth

Hypodermically also in spasmodic asthma, in narcotic poisoning, in aiding the reduction of hernia, and, with Strychnine, in lessening the craving for Alcohol. *See* also Atropine and Belladonna

In morphinism —*B M J E* '94, 1 20

In hernia, the hypodermic injection of $\frac{1}{10}$ grain of Atropine was followed by immediate spontaneous reduction. In four subsequent cases $\frac{1}{10}$ to $\frac{1}{20}$ grain were used. In another case, a second injection necessary, and in a sixth case three injections —*B M J E* '02, 11 92

In intestinal obstruction, three injections of $\frac{1}{10}$ grain each —*B. M. J. E.* '01, 11 48, *M. A.* '02, 362

In asthma, $\frac{1}{2}$ milligramme internally, increasing dose every second or third day by $\frac{1}{2}$ milligramme until patient is taking 4 milligrammes a day, gradually diminishing doses after a time. Duration of treatment, four to six weeks. Second and third course recommended at interval of six months —*Fr.* lxii 698

In broncho-pneumonia in children — *P*, lxx 698

Warning against the indiscriminate use of Atropine in eye diseases, particularly in the elderly — *B M J* '02, i 267

In 192, and '07, ii 951 are the details of the treatment for inebriety

Diabetes successfully treated — *B M J* '07, ii 28

Morphinomania successfully treated by Atropine and Strychnine — *B M J* '07, i 1173

In the treatment of infantile syphilitic iritis (*P*, lxxv 563), the local use of a solution of 2 grains of Atropine Sulphate to the oz of Distilled Water, together with theunction of mercurial ointment, is recommended

2 or 3 drops of a 4 grains to the oz solution instilled into the eye daily, or where the patient objected one eye was merely bandaged, thus altering the principal focus the motion of objects is not so uniformly transmitted to the brain and tendency to sea-sickness is diminished — *B M J* '05, i 1090

In the treatment of inebriety (*B M J* '05, ii 1691) the hypodermic injection of Atropine combined with Strychnine has proved of great value in relieving the intensity of the appetite, and when thoroughly pushed it confers indifference to Alcohol $\frac{1}{100}$ grain Atropine increased to $\frac{1}{80}$ grain with $\frac{1}{10}$ grain Strychnine; also, the injection of 1 minim of Liquor Atropine Sulphatis, with 4 minims of Liquor Strychnine Hydrochloridi, twice daily for a month, then once daily for a fortnight, then every second day for a month — *B M J* '06, i 515

Drops of Atropine Sulphate, 1 grain, Cocaine Hydrochloride, 1 grain, Adrenalin solution (1 in 1000), 30 minims, Water, 24 dim, applied 1 drop in the eye every 3 hours in acute conjunctivitis — *M P*, '05, ii 308

Two cases of toxic symptoms following the use of Atropine drops to the eyes of children In one case drops ' ' ' and in the other 1 grain of Atropine to the oz were used — *L* '06, ii 904

Dose. — $\frac{1}{200}$ to $\frac{1}{100}$ grain = 0.0003 to 0.0006 gramme.

Dan, Dutch, Fr, Ger, Ital, Norw, Russ and Swiss give the maximum single dose as 0.001 gramme, Ger, Ital and Russ, maximum daily dose 0.003 gramme, Fr, 0.002 gramme

Prescribing Notes — *The Sulphate is best adapted for aqueous Solutions, and the pure Alkaloid for Ointments. Can be given in pill and triturated with Milk Sugar and mixed with Diluted Glucose. Generally given in solution.*

Official Preparations — Lamellæ Atropinae, and Liquor Atropinae Sulphatis

Not Official — Glycerinum Atropinae, Guttæ Atropinae Sulphatis, Guttæ Atropinae cum Cocaina, Injectio Atropinae Hypodermica, Linimentum Atropinae, Linimentum Atropinae et Chloroformi, Pilulæ Atropinae et Morphinae, Pessus Atropinae, Pilulæ Atropinae, Atropinae Methylbromidum, Atropinae Salicylas, Atropinae Valerianas, Euphthalmine Hydrochloridum, Guttæ Euphthalmine Hydrochloridi, Lamellæ Euphthalmine

Atropine is used as an antidote in poisoning by Phosphorus, Morphine, Aconite, Gelsemine, Hydrocyanic Acid, Muscarine, Nerve poisons, and Pilocarpine

Antidotes — In case of poisoning by Atropine, the antidotes are the same as for Belladonna, *q v*

Foreign Pharmacopœias — Official in Aust, Belg, Dan, Dutch, Fr., Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U.S

Tests — The distinguishing tests for Atropine Sulphate are the melting point, which should be 189° C (347.2° F), the mydriatic action produced even by highly diluted solutions of the salt, the production of a purple-violet coloration when a small quantity is moistened with Nitric Acid evaporated on a water-bath and the residue treated with an Alcoholic Potassium Hydroxide Solution; the production of a white precipitate, on the addition of Sodium Carbonate to a saturated aqueous solution, and when this precipitate is collected

it should answer the tests described under Atropine. The aqueous solution yields on acidification with diluted Hydrochloric Acid and addition of Barium Chloride Solution, a white precipitate insoluble in Hydrochloric Acid.

The melting point given in the *BP* and *PG* is open to criticism (*PJ* '98, ii 195). Will gives 196°C ($384\ 8^{\circ}\text{F}$), *USP* about $189\ 9^{\circ}\text{C}$ ($373\ 5^{\circ}\text{F}$), and when free from Hyoscyamine about 188°C ($370\ 4^{\circ}\text{F}$), Hesse, 180° to 181°C (356° to $357\ 8^{\circ}\text{F}$), Merck, 189° to 191°C ($372\ 2^{\circ}$ to $375\ 8^{\circ}\text{F}$), whilst a salt prepared by Jowett from pure Atropine melted at 190°C (374°F). 3 c c of a 1 in 60 aqueous solution mixed with 1 c c of Sodium Hydroxide Solution (15 p c) yields a precipitate, but a solution of the same strength is unaffected by a corresponding amount of Ammonia Solution. A pleasant aromatic odour is evolved when a small quantity of the salt, which has been heated until white fumes are disengaged, is warmed with Sulphuric Acid until it commences to turn brown, and then a small quantity of Water is carefully added, the addition of a few crystals of Potassium Permanganate produces an odour of Essential Oil of Bitter Almonds.

The more generally occurring impurities are Sulphates of mydriatic alkaloids other than Atropine, *eg*, Hyoscyamine and Scopolamine, readily oxidisable organic impurities, Morphine and mineral matter. Hyoscyamine and Scopolamine Sulphates are indicated by their optical activity, organic impurities by the colour imparted to a solution of the salt in concentrated Sulphuric Acid, Morphine by the red coloration produced on the addition of Sulphuric Acid followed by Nitric Acid to a little of the salt, and mineral matter by the ash left on ignition which should be nil. Atropine Sulphate contains 85.5 p c of Atropine and 14.5 p c of Sulphuric Acid.

Sulphuric Acid—If 1.5 c c Sulphuric Acid be added to 0.01 gramme of Atropine Sulphate which has been heated in a test-tube until the evolution of white vapours occurs, and then warmed until the mixture begins to turn brown, then on immediately and carefully adding Water to this, a pleasant characteristic aromatic odour comes off. After the addition of a crystal of Potassium Permanganate the liquid smells of Essential Oil of Almond, *PG*. See also *USP*, under Atropine.

Vitali's Test—0.01 gramme Atropine Sulphate heated to dryness in a porcelain dish on a water-bath with 5 drops fuming Nitric Acid, leaves a faintly yellow residue, which on pouring over it an Alcoholic Solution of Potassium Hydroxide, and warming, assumes a violet colour, *PG*.

Preparations

LAMELLÆ ATROPINÆ. DISCS OF ATROPINE

Discs of Gelatin, each weighing about $\frac{1}{10}$ gram (1.3 milligrammes) and containing $\frac{1}{1000}$ gram (0.013 milligramme) of Atropine Sulphate.

Gelatin Discs, each containing 0.001 gramme Atropine Sulphate, are official in Sweden. It contains $\frac{1}{10}$ milligramme.

LIQUOR ATROPINÆ SULPHATIS. SOLUTION OF ATROPINÆ SULPHATE.

Atropine Sulphate, 1, Salicylic Acid, $\frac{1}{2}$, Distilled Water, 100
(1 in 100)

Dose.— $\frac{1}{2}$ to 1 minim = $\frac{1}{20}$ to $\frac{1}{10}$ grain of Atropine Sulphate.

Foreign Pharmacopœias—Official in Port, 1 in 100. Not in the others.

Not Official.

GLYCERINUM ATROPINÆ—Atropine Sulphate, 25 $\frac{1}{2}$ grains, dissolved in Water, 5 fl oz., add Compound Tincture of Lavender, 100 minims, and Glycerin, to 20 fl oz. 100 c c contains 0.25 gramme Atropine—*St Thomas's*

Atropine Sulphate, 1 $\frac{1}{2}$ grains, Water, 2 dim., Compound Tincture of Lavender, 5 minims, Glycerin, to 1 oz.—*University*

Atropine Sulphate, 0.25, Distilled Water, 25, Compound Tincture of Lavender, 1, Glycerin, q s to produce 100—*B P C*

GUTTÆ ATROPINÆ SULPHATIS—Atropine Sulphate, 1, 2 or 4 grains, Di-tilled Water, 1 oz.—*London Ophthalmic*

GUTTÆ ATROPINÆ CUM COCAINA—Atropine Sulphate, 2 grains; Cocaine Hydrochloride, 10 grains, Distilled Water, 1 oz.—*London Ophthalmic.*

INJECTIO ATROPINÆ HYPODERMICA.—Atropine Sulphate, 2 grains, Water, 1 oz

Dose—2 to 4 minims = $\frac{1}{20}$ to $\frac{1}{10}$ grain of Atropine Sulphate.

B P C Injection contains 0.12 p c

INJECTIO ATROPINÆ ET MORPHINÆ HYPODERMICA. See *MORPHINÆ ACETAS*

LINIMENTUM ATROPINÆ—Atropine Sulphate, 88 $\frac{1}{2}$ grains; Compound Tincture of Lavender, 100 minims, Alcohol (90 p c), to 20 fl oz.—*St Thomas's.*

This has been incorporated in the *B P C* as follows—

Atropine Sulphate, 0.40, Compound Tincture of Lavender, 1, Alcohol, q. s. to produce 100

LINIMENTUM ATROPINÆ ET CHLOROFORMI—Atropine Liniment 5, Chloroform, 1—*St Thomas's*

This has been modified in the *B P C* as follows—

Chloroform, 12.50, Atropine Liniment, q s to make 100

PILULA ATROPINÆ—Atropine Sulphate, $\frac{1}{16}$, $\frac{1}{8}$, $\frac{1}{4}$, $\frac{1}{2}$ grain; Liquorice Powder, 2 grains, Tragacanth Powder, 1 grain, Mucilage of Acacia, q s—*Brompton*

PILULA ATROPINÆ ET MORPHINÆ—Atropine Sulphate, $\frac{1}{16}$ grain, Morphine Hydrochloride, $\frac{1}{4}$ grain, Milk Sugar, 1 grain—*St Thomas's.*

This has been incorporated in the *B P C*

PESSUS ATROPINÆ—Atropine, $\frac{1}{2}$ grain, Conine, 1 minim; Oil of Theobroma, to 120 grains—*Samaritan*

ATROPINÆ METHYLBROMIDUM (Mydrasine)—A white crystalline powder, readily soluble 1 in 1 of Water, soluble 1 in 10 of Alcohol (90 p c), insoluble in Ether. Employed in the form of a 1 to 2 p c solution either plain, or as a solution containing 1 p c of Cocaine. A valuable succedaneum for Atropine Sulphate. Free from its disadvantages and possessing its advantages. Invaluable in determining static refraction or in any ophthalmoscopic examination, where dilatation of the pupil is indispensable. In large quantities (1 to 2 p c solution) it has the same action on the pupil and accommodation as Atropine Sulphate. In moderate quantities (1 drop of 1 p c solution) the mydrasis produced may last about twenty-four hours, but the paralysis of accommodation disappears in a few hours. In small quantities (1 drop 0.5 p.c. solution containing 1 p c of Cocaine), considerable dilatation of the pupil results with scarcely any appreciable paresis of accommodation. Atropine Methylbromide is thus a valuable diagnostic agent in the beginning of iritis, but should there be no iritis, the inconvenience produced is less than if Atropine had been instilled. One drop of a 1 p c solution produces a maximum dilatation of the pupil in from 30 to 45 minutes. Cocaine intensifies the dilatation; Eserine quickly diminishes the pupillary dilatation produced by Methylbromide.—

B M J E '03, 11 52 Has been used in doses of $\frac{1}{10}$ to $\frac{1}{2}$ of a grain = 0.0065 to 0.013 gramme in pill form in the treatment of the night sweats of phthisis. In addition to its mydriatic effect it has a distinct sedative and analgesic effect (*B M J E* '06, 1 72), for ophthalmic purposes it may be used in strengths from $\frac{1}{4}$ to 5 p c. A useful average strength is 1.0 p c.

Tests.—Atropine Methylbromide has a melting point of 222° to 223° C (431.6° to 433.4° F), its aqueous solution yields a whitish precipitate with Mayer's reagent, the residue remaining after evaporating a small quantity of the salt with a few drops of concentrated Nitric Acid yields a purple violet coloration when treated with Alcoholic Potassium Hydroxide Solution, its aqueous solution yields a white curdy precipitate with Silver Nitrate solution and a yellow coloration with Chlorine Water, the latter when shaken with Chloroform yielding a yellowish brown solution. It should leave no residue when ignited with free access of air.

ATROPINÆ SALICYLAS, $C_{17}H_{13}NO_3 \cdot C_7H_5O_2$, eq. 424.06.—A white crystalline powder, only slightly soluble in Water. Introduced as a substitute for the Sulphate in ophthalmic practice, but its aqueous solution does not keep so well as that of the latter. The author prepared 1 p c solutions of each salt, and the Salicylate developed a growth more quickly than the Sulphate. To make the solution keep well an excess of Salicylic Acid is required, and then it is irritating to the eye.

Tests.—The aqueous solution yields with Mayer's reagent a whitish precipitate, a crystal evaporated on a water bath with concentrated Nitric Acid leaves a residue which when moistened with a few drops of Alcoholic Potassium Hydroxide Solution yields a purplish violet coloration. The diluted aqueous solution yields with Ferric Chloride Test solution a deep violet coloration. It should leave no residue when ignited with free access of air.

It should contain 67.7 p c of Atropine and 32.3 p c of Salicylic Acid.

Liquor Atropinæ Salicylatis—Atropine, 5 grains, Salicylic Acid, 7½ grains, Water, 10 oz.—*Charing Cross*

ATROPINÆ VALERIANAS, $C_{17}H_{13}NO_3 \cdot C_8H_{10}O \cdot H_2O$, eq. 406.24.—Colourless, or white hygroscopic rhomboid crystals, having an odour of Valerianic Acid, and becoming coloured on exposure to light. Readily soluble in Water and in Alcohol (90 p c). Antispasmodic, antineuralgic. Recommended for internal administration.

Dose.— $\frac{1}{8}$ grain = 0.001 gramme. **Official in Mex. and Port.**

Tests.—The salt softens at 20° C (68° F), and melts at 32° C (89.6° F). The aqueous solution when acidified with a mineral acid throws out an oily fluid which collects on the surface of the liquid, Sodium or Potassium Hydroxide Solution produces a white precipitate. A crystal with a few drops of concentrated Nitric Acid evaporated to dryness yields a residue which assumes a purplish-violet coloration when moistened with a few drops of Alcoholic Potassium Hydroxide Solution. When ignited with free access of air it should leave no residue. The crystallised salt should contain 70.6 p c of Atropine and 24.9 p c of Valerianic Acid.

EUPHTHALMINÆ HYDROCHLORIDUM (Phenylglycoyl-*n*-methyl β vinyl diacetone alkamine hydrochloride).—White crystalline powder. Readily soluble in Water. Mydriatic. Introduced as a substitute for Atropine and Homatropine, and used as a 2 to 5 p c solution. Stated to weaken the accommodation only to a very slight extent, but has no appreciable effect on the conjunctival vessels or on the corneal epithelium, and causes no hyperæmia, and effects soon pass off.—*B M J* '99, 11 775, *L* '99, 11 458, *P* lxiv 476. The free base Euphthalmine crystallises in six sided prisms. A readily soluble Euphthalmine Salicylate has also been prepared.

Guttæ Euphthalminæ Hydrochloridi—Euphthalmine Hydrochloride, 10 grains, Distilled Water, 1 oz.—*London Ophthalmic*

Lamellæ Euphthalminæ.—Each disc contains $\frac{1}{2}$ grain Euphthalmine.—*London Ophthalmic*

AURANTII CORTEX.

FR, BIGARADIER, GER, POMERANZENSCHALE, ITAL, ARANCIO AMARO;
SPAN, NARANJA AMARA

Both the fresh and the dried outer part of the Pericarp of *Citrus Aurantium*, var *Bigaradia*, are official

In India and the Eastern Colonies, Auranti Cortex Indicum (*Ind* and *Col Add*) may be used. It is the dried portions of the varieties of *Citrus Aurantium* grown in India and Ceylon

Medicinal Properties—Carminative and bitter stomachic. The Tincture and Syrup are largely used as *fructus* agents.

Prescribing Note—If Orange Peel should not be prescribed with Tincture of Peppermint, the latter would be blackened.

Official Preparations—Of the **Fresh Peel**, Tinctura Aurantii and Vinum Aurantii. Of the **Tincture**, Syrupus Aurantii, contained in Tinctura Quininae, Syrupus Aromaticus and Syrupus Cascariae Aromaticus. Of the **Dried Peel**, Infusum Aurantii and Infusum Aurantii Compositum, used in the preparation of Infusum Gentianae Compositum, Spiritus Amomaci Compositus, Linctura Cichorii Composita, and Tinctura Gentianae Composita.

Not Official—Oleum Aurantii Corticis, Elixir Aurantii, Elixir Aromaticum, Elixir Simplex, Infusum Aurantii Concentratum, Infusum Aurantii Compositum Concentratum, Spiritus Aurantii Compositus, Vinum Aurantii Detannatum.

AURANTII CORTEX RECENS. FRESH BITTER-ORANGE PEEL

The Fresh outer part of the Pericarp of *Citrus Aurantium*, var *Bigaradia*, Hook f

Foreign Pharmacopœias—Official in Belg, Fr, Mex, Port and Span, U S, *Citrus Aurantium*. The following use the unripe fruit: Ger, Jap, Norw, Russ, Swed and U S

Descriptive Notes.—The fresh rind of the Seville or bitter orange *Citrus Aurantium*, L, var *Bigaradia*, Hook f, is official. It is characterised by its reddish or deep orange red colour and its rough glandular surface. It should retain but little of the white spongy portion. The taste is bitter and aromatic. It is most easily purchased in the fresh state in February and March, when it usually arrives in this country.

Preparations

SYRUPUS AROMATICUS. AROMATIC SYRUP

Tincture of Orange, 1, Cinnamon Water, 1; Syrup, 2. The turbid fluid, formed by mixing the Tincture and Cinnamon Water, is cleared by filtration through Tale, before the Syrup is added.

Dose.— $\frac{1}{2}$ to 1 fl drm. = 1.8 to 3.6 cc

SYRUPUS AURANTII. SYRUP OF ORANGE.

Tincture of Orange, 1, Syrup, 7

(1 in 8)

Dose.— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Foreign Pharmacopœias—Official in Hung, Peel, weak Spirit, Sugar, and Tincture, Belg, Fluid Extract 1, Syrup 19, Dutch, Peel, Water, and Sugar,

Dan, Fr, Ital and Swed, Peel, Spirit, Water, and Sugar, Austr, Ger, Russ and Swiss, Peel, Wine, and Sugar, Jap, Tincture of Bitter-Orange Peel 3, Syrup 17, Norw, Tincture 1, Syrup 9, Mex, Alcoholatura 1, Syrup 9, Port, Peel, Boiling Water and Sugar, Span (Jarabe de Corteza de Naranja), Water of Orange Peel and Sugar, (Jarabe de Corteza de Naranja Amarga), Wine of Bitter Orange Peel and Sugar All by weight, except U S

Syrupus Aurantii (U S)—Tincture of Sweet Orange 5, is triturated with Magnesium Carbonate 1, and Water 40, gradually added, filtered, sufficient Water added to produce 45, Citric Acid 0.5 dissolved in the filtrate, then Sugar 82, and sufficient Water added to produce 100 by volume

TINCTURA AURANTII TINCTURE OF ORANGE

Macerate 1 of Fresh Bitter Orange Peel, cut small, with 4 of Alcohol (90 p c) (1 in 4)

Formerly called Tinctura Aurantii Recentis, and 6 of Fresh Peel made 20 of Tincture

Dose —30 to 60 minims = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Belg, Fresh Peel 1, Alcohol (60 p c) 5, Dutch, Peel 1, Alcohol (70 p c) 5, Fr (Alcoolature d'Orange), Fresh Peel 1, Alcohol 2, Ital, Peel 1, Alcohol 2, both by weight U S (Tinctura Aurantii Dulcis), from Fresh Peel, 1 in 5, not in the others The following are made with Dried Peel Austr, Dan, Dutch, Fr, Ger, Hung, Jap, Norw, Russ, Swed, Swiss and U S, 1 and 5, all by weight, except U S Not in Port

Tests—Tincture of Orange has a sp gr of about 0.880, contains about 2 p c w/v of total solids and about 74 p c w/v of Absolute Alcohol

VINUM AURANTII ORANGE WINE

A sherry-coloured weak alcoholic liquid, prepared by the fermentation of a saccharine solution containing Fresh Bitter Orange Peel

Foreign Pharmacopœias—Official in Span, Dried Bitter Orange Peel 1, Carinena or Alicante Wine 20 Belg and Swiss have a compound wine, but they vary considerably in composition

The Orange Wine of commerce

Tests.—Orange Wine has a sp gr of about 1.030, it contains about 11.5 p c of total solids It is officially required to contain 10 to 12 p c w/v of Ethyl Hydroxide Good commercial samples contain from 15 to 18 p c w/v of Absolute Alcohol In testing for Salicylic Acid 1 fl oz of the Wine is rendered alkaline by the addition of Sodium Hydroxide Solution, and shaken with Ether, the ethereal solution separated and rejected, the Wine may then be rendered acid with diluted Sulphuric Acid and again shaken with Ether, the ethereal solution separated, carefully washed with Water till free from mineral acid and shaken with Water to which a drop or two of Ferric Chloride T S has been added, Salicylic Acid, if present, is indicated by the violet coloration produced The B P performs the test for Salicylic Acid on the distillate obtained from a mixture of 50 c c of Wine, 50 c c of Water and 5 c c of Normal Volumetric Sulphuric Acid Solution, rejecting the first 10 c c distilled and shaking the balance with Ether Ferric Chloride T S is used as a reagent, and the test is carried out on the residue left on evaporation

Not Official

INFUSUM AURANTII CONCENTRATUM —Dried Bitter Orange Peel, in No 10 powder, 40, Tincture of Orange, 5, Alcohol (90 p c), 22 5 Dilute Chloroform Water (1 in 1000), *q s* to make 100 Prepare by repercolation Dose — $\frac{1}{2}$ to 1 fl dm —*Farr and Wright, P J '06, 1 165 and '07, 1 621, C D '06, 1 252, Y B P 1907, 250*

This appears in the *B P C*

INFUSUM AURANTII COMPOSITUM CONCENTRATUM —Dried Bitter-Orange Peel, in No 10 powder, 20, Dried Lemon Peel, in No 10 powder, 5, Cloves, freshly powdered, 2 5, Tincture of Lemon, 5, Tincture of Orange, 5, Alcohol (90 p c), *q s* Dilute Chloroform Water (1 in 1000), *q s* to make 100 Macerate the powdered Cloves in 20 of the Alcohol for 12 hours, filter through Cotton-Wool, and pass through the same sufficient Alcohol to make the filtrate measure 20, add the tinctures and set aside, mix the other powders, and submit them to macero-expression with dilute Chloroform Water, adding the mixed tinctures to the reserved portion Dose — $\frac{1}{2}$ to 1 fl dm —*Farr and Wright, P J '06, 1 165 and '07, 1 621, C D '06, 1 252, Y B P 1907, 249*

This appears in the *B P C*

VINUM AURANTII DETANNATUM —Orange Wine, 1 gallon, Gelatin, in No 100 powder, $\frac{1}{2}$ oz., macerate for 24 hours and decant

It is recommended by *F C J* Bud that Gelatin in No 100 powder should be used in place of Gelatin cut small, as recommended in the *B P C Formulary* 1901, it being found possible by this process to completely detannate an average sample of Wine in 24 hours with the aid of occasional shaking, or in 8 hours if frequently agitated Care must be taken to keep the temperature of maceration at or below 15 5° C, or during extremely hot weather the Gelatin will probably pass into solution —*P J '99, 1 183*

This suggestion has been incorporated in the *B P C* as follows —

Orange Wine 100, Gelatin, in No 100 powder, 0 15 Macerate for 24 hours at a temperature not exceeding 15 5° C, with frequent agitation, and afterwards decant.

OLEUM AURANTII CORTICIS —A volatile Oil, extracted by mechanical means from Fresh Orange Peel, both varieties of Orange Peel are used, that from *Citrus Aurantium*, L, var *Bigaradia*, Hook, is known as *Essence de Bigarade*, and that from *Citrus Aurantium*, L, as *Essence de Portugal*, the former yields the finest Oil

A pale yellowish liquid, with neutral reaction, having the odour of Orange Peel At least 90 p c of the Oil consists of dextro-rotatory Limonene

By keeping, the Oil becomes thicker and acquires a disagreeable terpeninathate taste, which may be prevented by mixing it while fresh with 10 p c of Absolute Alcohol It should be kept in well-stoppered dark amber-tinted glass bottles in a cool place

Solubility —Soluble 1 in 7 of Alcohol (90 p c), and in all proportions of Absolute Alcohol

Foreign Pharmacopœias —Official in Austr and Belg., sp gr 0 848 to 0 852, Dutch, sp gr 0 850 to 0 870, Fr, sp gr 0 848 to 0 853, Hung. and Jap, sp gr 0 850 to 0 860, Port, sp gr 0 835 to 0 850, Mex., sp gr 0 837; Spain, 0 835 to 0 844, U S, sp gr 0 842 to 0 846 at 25° C (77° F.), not in Dan, Ger, Ital, Norw, Russ, Swed or Swiss.

Tests —The oil of sweet orange has a sp. gr of 0 848 to 0 852, that from bitter orange 0 854 to 0 857 Both have a characteristic orange odour The oil has a boiling point of 175° to 180° C (347° to 356° F), between which temperatures about nine-tenths of the oil distils over The oil of sweet orange has an optical rotation in a tube of 100 mm diameter of +96° to +98° at 20° C (68° F), that of the bitter orange is between +90° and +98°

The more generally occurring sophistications are Turpentine Oil and the terpenes remaining from the manufacture of terpeness Oil of Lemon and Oil of Orange Turpentine Oil may generally be detected directly by the optical rotation, or by a determination of the optical activity of the first 10 p c fraction,

The optical rotation of this first 10 p c fraction should not materially differ from that of the original oil

The terpenes remaining from the manufacture of terpeneless Lemon Oil are readily detected by the optical rotation, but terpenes from terpeneless Oil of Orange are detected with much greater difficulty. They reduce the colour of the oil and produce a difference in the odour and taste

ELIXIR AURANTII (formerly *U S*, now omitted)—Sprinkle or spray 1 fl oz of Oil of Orange over 2 oz of Cotton-Wool, pack it tightly in a percolator and pass through it a mixture (Alcohol 1, Water 3), sp gr 0.971, till 200 fl oz of a clear percolate are obtained, in which dissolve, without heat, Sugar 100 oz, all by weight

A better method of disseminating the Oil is to sprinkle it upon blotting paper pulp this with the diluted Alcohol, allow it to stand for 24 hours, and filter

ELIXIR AROMATICUM—Compound Spirit of Orange, 1 2, Syrup, 87 5, Purified Talc, 3, Alcohol (95 p c), *q s*, Distilled Water, *q s* to produce 100. To the Compound Spirit of Orange add enough Alcohol to make 25, to this solution add the syrup in several portions, agitating after each addition, then 87.5 of Distilled Water. Mix the Purified Talc with the liquid and then filter through a wetted filter, returning the first portions of the filtrate until a transparent liquid is obtained. Lastly, wash the filter with a mixture of Alcohol 1 volume and Distilled Water 3 volumes, until the product measures 100—*U S P*

This has been incorporated in the *B P C*, which appears to direct twice as much Compound Spirit of Orange, but the *B P C* Compound Spirit is only half the strength of the *U S P*. This, however, has been altered in the *B P C Supplement*, the Spiritus Aurantii Compositus (see below) has now been doubled in strength, and half the quantity of it used, and the amended formula for Elixir Aromaticum will agree with that given above for *U S P*

ELIXIR SIMPLEX—Oil of Bitter Orange, 30 minims, Alcohol (90 p c), 6 fl oz, dissolve and add Distilled Cinnamon Water, 7 fl oz, Syrup, 7 fl oz, mix. Filter through paper moistened with Alcohol (45 p c) and well sprinkled with kaolin, returning the first portions of filtrate until it passes through bright—*B P C Formulary* 1894 omitted in 1901

Dose—20 to 60 minims = 1 2 to 3 6 c c

Tincture of Orange, 7 50, Syrup, 40, Distilled Water, *q s* to make 100.—*B P C*

This is the same formula as given under Pepsin

SPIRITUS AURANTII COMPOSITUS (*U S*)—Oil of Orange, 10, Oil of Lemon, 2½, Oil of Coriander, 1, Oil of Anise, ½, Alcohol (95 p c), to make 50

The *B P C* Spiritus Aurantii Compositus employ Alcohol (90 p c) to make 100 and is therefore half this strength. This has been altered in the *B P C Supplement*, reducing the quantity of Alcohol (90 p c) to make 50, so that the formula will correspond in strength to the *U S* preparation, but using Alcohol (90 p c) in place of *U S P* Alcohol (95 p c)

AURANTII CORTEX SICCATUS. DRIED BITTER-ORANGE PEEL

The dried outer part of the Pericarp of *Citrus Aurantium*, L., var *Bigaradia*, Hook f

Official Preparations—Infusum Aurantii and Infusum Aurantii Compositum

Foreign Pharmacopœias—Official in Austr, Dan, Dutch, Fr, Ger, Hung, Ital (Arancio Amaro), Jap, Norw, Port. (Laranjeira Azeda), Russ, Swed and Swiss, *U S*, Aurantii Amari Cortex, also Aurantii Dulcis Cortex.

Descriptive Notes.—Bitter-Orange Peel occurs in English commerce in three forms, viz, English, exotic and Maltese. The English is more carefully dried, externally and nearly white on the inner surface, in long strips about $\frac{1}{8}$ in (8 mm.) wide and more or less curled in drying. The exotic is less carefully dried and of a duller orange-red or brownish-red colour externally, and a dirty white colour on the inner surface. The Maltese resembles the exotic, except that it occurs in very slender strips about 2 lines wide only, is cut up into shorter pieces, and has less of the mesocarp or white spongy portion attached to the zest or rind. The official description simply directs that it should be in thin strips, but as the colour must be orange-red, it is evident that the English-dried kind is intended. The *U S P* describes the dried peel as of a brownish-green colour, and in strips or quarters, thus apparently admitting the badly dried W Indian peel which is usually dried in quarters. The *P G* describes the dried peel as brownish, and directs that it should be prepared by softening the rind in cold Water for a quarter of an hour, pouring off the Water and keeping the peel in a cool place until the next day when the white spongy tissue should be cut off and the outer portion dried. Dried Orange Peel if long kept loses its bright colour and becomes brownish-red.

Preparations

INFUSUM AURANTII. INFUSION OF ORANGE PEEL.

Dried Bitter-Orange Peel, 1, boiling Distilled Water, 20. Infuse for 15 minutes (1 in 20)

Dose $-\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc

Fr (Tisane d'Oranger), Leaves, 5, boiling Water, 1000

INFUSUM AURANTII COMPOSITUM. COMPOUND INFUSION OF ORANGE PEEL.

Dried Bitter-Orange Peel, $\frac{1}{2}$ oz, Fresh Lemon Peel, $\frac{1}{2}$ oz, Bruised Cloves, 55 grains, boiling Distilled Water, 20 oz. Infuse for 15 minutes (1 in 40)

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc.

TINCTURA AURANTII. See AURANTII CORTEX RECENS

Formerly two Tinctures were official, one from Fresh Peel and the other from Dried Peel, the latter is now omitted

AURANTII FLORIS AQUA.

ORANGE-FLOWER WATER

NO Syn—AQUA NAPHE

Fr, L'EAU DE FLEUR D'ORANGER; GER, POMERANSENBLÜTENWASSER; ITAL, ACQUADI FIORI DI ARANCIO; SPAN., AGUA DESTILADA DE AZAHAR

Commercial Orange-flower Water prepared by a process of distillation from the Flowers of the Bitter-Orange tree, *Citrus*

Aurantium, L, var *Biganadra*, Hook f, diluted with twice its volume of Distilled Water. It keeps best in the undiluted state, and should therefore be diluted only as required.

U S directs the Triple Extract to be diluted with an equal volume of Distilled Water. Swiss use the undiluted Water.

Medicinal Properties—Both the Water and the Syrup are used as flavouring agents, about 1 of the Concentrated Water to 8 of Distilled Water, it is also used in eye lotions.

Official Preparation—Syrupus Aurantii Floris. Contained in Mistura Olei Ricini, and Syrupus Calcii Lactophosphatis.

Not Official—Oleum Aurantii Florum (Oleum Neroli).

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr (Eau Distillée de Fleur d'Oranger), Hung, Ital (Acqua Distillata di Arancio) Jap, Mex (Agua destilada de corteza de naranja amarga), Port (Água de Flores de Laranjeira), Russ, Span (Agua Destilada de Azahar), Swed, Swiss and U S. Not in Dan, Ger, or Norw.

Preparation

SYRUPUS AURANTII FLORIS SYRUP OF ORANGE FLOWER

Dissolve 6 of Refined Sugar in 2 of boiling Distilled Water, add 1 of undiluted Orange-flower Water of commerce, and make up the total weight to 9 with recently boiled Distilled Water.

Dose— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Foreign Pharmacopœias—Official in Austr, O F W 1, Sugar $1\frac{1}{2}$, Belg, O F Spirit (1 Oil in 100) 1, Syrup 199, Fr and Mex, O F W 10, Sugar 18, Port, O F W 7, Sugar 13, Span and Swiss, O F W 36, Sugar 64, all by weight. U S, Sugar 85, O F W to measure 100. Not in the others.

Not Official

OLEUM AURANTII FLORUM *Syn* OLEUM NEROLI—A volatile Oil, obtained by distilling fresh Orange flowers with Water. The watery distillate constitutes the Aqua Floris Aurantii Conc of commerce. The finest Oil is obtained from the Bitter-Orange, that from the Portugal or Sweet-Orange is not so good. From the leaves, twigs and immature fruits of both varieties is obtained the commercial Oil of Petit Grain.

A yellowish or brownish limpid liquid, with neutral reaction, having a powerful odour of Orange flowers.

Solubility—Soluble in all proportions of Alcohol (90 p c) or in Absolute Alcohol.

Foreign Pharmacopœias—Official in Austr, Belg and Swiss, sp gr 0.870 to 0.880, Mex, sp gr 0.870 to 0.878, Fr, sp gr 0.875 to 0.880, Span, sp gr 0.850 to 0.900, Ital, sp gr 0.872 to 0.890, Jap, sp gr 0.86 to 0.88, Port, sp gr 0.874 to 0.878. Not in Dan, Dutch, Ger, Hung, Norw, Russ, Swed or U S.

Tests—The Oil has a specific gravity of 0.870 to 0.890 and a powerful characteristic odour of Orange flowers. It should possess an optical rotation in a tube of 100 mm diameter of $+2^{\circ}$ to $+5^{\circ}$ and occasionally as high as $+8^{\circ}$. It should form a clear solution in $1\frac{1}{2}$ to 2 volumes of Alcohol (80 p c). A determination of the Saponification value affords a useful criterion of the purity of an Oil, that of genuine Oils being between 20 and 52 corresponding to 7 to 18 p c of Esters calculated as Linallyl Acetate.

The more generally occurring sophistications are Bergamot and Petit Grain Oils, which being composed largely of the same chemical constituents as Neroli

Oil can only be detected when in relatively large amounts being next to impossible. The value here affords the recognition of the best criterion.

Not Official
AURI BROMIDUM

AuBr₃, eq 433.75

In dark brown masses, soluble in Water. It has been used on the Continent for the relief of hysteria and epilepsy.

The salt should be kept in small bottles of a dark amber tint.

The Tribromide obtained from it is soluble about 1 in 75 of Water.

It appears to be about ten times more active than the more commonly used Bromides, and has been given in $\frac{1}{2}$ (increased to $\frac{1}{3}$) grain doses in severe cases of hysteria and epilepsy — *L* '90, i 869.

Dose — $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.0067 to 0.016 gramme.

Prescribing Notes — Dispensed in pills with *Massa Kauloni*, or in compressed Tablets.

Official in Mex., Bromuro de Oro.

AURI ET POTASSII BROMIDUM — Brownish-black needle-shaped crystals. Readily soluble in Water. Used for the same purposes as for the Tribromide.

Dose — $\frac{1}{2}$ grain = 0.021 gramme.

LIQUOR AURI ET ARSENI BROMIDI — Arsenious Acid, 0.25 gramme, Gold Tribromide, 0.325 gramme, Bromine Water and Distilled Water, of each a sufficient quantity to make 100 c.c. — *U.S.N.F.* 1896 and 1906.

10 minims of this solution contain $\frac{1}{2}$ grain of Tribromide of Gold and the equivalent of $\frac{1}{8}$ grain of Tribromide of Arsenic.

Arsenious Acid, in powder, 40 grains, Potassium Carbonate, 40 grains, Bromine, 100 grains, Gold, in leaf, 13.5, Distilled Water, q.s. to 20 fl. oz. (Wright) — *Y.P.B.* 1896, 354.

10 minims contains an amount of Arsenium in combination equal to $\frac{1}{8}$ grain of Arsenious Acid and $\frac{1}{2}$ grain of Gold Tribromide.

This has been incorporated in the *B.P.C.* under the title **Liquor Auri et Arsenii Bromatus**.

This liquor has been found useful in rheumatism — *L* '95, ii 921, '00, i 72; it is also combined with Mercury Oxybromide in syphilitic affections.

Not Official.
AURI CHLORIDUM.

Under this heading are arranged the following varieties —

- 1 **Pure Chloride of Gold**, AuCl₃, containing about 65 p.c. of metallic Gold. Official in Port. (Chloreto de Ouro), and Mex. (Cloruro de Oro).
- 2 **Chloride of Gold and Sodium** (Commercial 'Chloride of Gold'), the crystallised double salt AuCl₃.NaCl.2H₂O, containing 50 p.c. of Gold. Official in Fr. (Chlorure d'Or et de Sodium), Port. (Chlorure de Ouro e de Sodio), and Mex. (Cloruro de Oro y Sodio).
- 3 **Commercial Chloride of Gold and Sodium**. Commercial Chloride of Gold and Sodium is the above crystallised salt mixed with an equal weight of Chloride of Sodium, and contains 25 p.c. of metallic Gold.
- 4 **Auri et Sodii Chloridum, U.S.** A mixture composed of equal parts of anhydrous Gold Chloride and anhydrous Sodium Chloride,

and which contains not less than 30 p.c. of metallic Gold
Official in Russ (Auro-natrium Chloratum)

The salts should be kept in well-stoppered bottles of a dark amber tint

Some foreign samples of commercial Chloride of Gold are the double Chloride of Gold and Potassium $\text{AuCl}_3 \cdot \text{KCl} 2\frac{1}{2}\text{H}_2\text{O}$, corresponding to about 47 p.c. of metal—*P J* (3) xxii 902

Medicinal Properties—It has been given on the Continent for amenorrhœa and secondary syphilis Chloride of Gold and Sodium has been used in tertiary syphilis, spinal sclerosis, hystero epilepsy, asthma, chorea, and in uterine affections

Dose— $\frac{1}{16}$ to $\frac{1}{2}$ grain = 0.004 to 0.016 gramme

Prescribing Notes—*It may be given in the form of pills made with Massa Kaolin, or in aqueous solution Its solutions should be protected from white light*

It is also used in photography

Not Official.

AZADIRACHTA INDICA

INDIAN AZADIRACH

Syns—NEEM BARK, MARGOSA BARK

The dried Bark of the stem of *Melia Azadirachta*, L. **Infusum Azadirachtæ Indicæ** (1 in about 109), dose $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 cc., and **Tinctura Azadirachtæ** (1 in 10), dose 30 to 60 minims = 1.8 to 3.6 cc., are official in the *Ind* and *Col Add* for India and the Eastern Colonies

BALSAMUM CANADENSE.

See TEREBINTHINA CANADENSIS

Not Official

BALSAMUM DIPTEROCARPI

GURJUN BALSAM

A balsamic exudation, obtained from the Trunk of *Dipterocarpus turbinatus*, Gærtn f., and other species by incision and the application of heat Imported from the East Indies It is an oleo resin, constituting a transparent liquid of the consistence of Olive Oil, lighter than Water, and of a dark brown sherry colour, slightly fluorescent

Medicinal Properties—Similar to those of Copaiba Useful for leprosy. Mr J D Hillis, of the Leper Asylum in British Guiana, is greatly in favour of it—*L* '80, 1 659, *MP* '89, 1 664, see also *L* '90, 1 136 Von Reissen gives Wood Oil internally, commencing with daily doses of 5 drops, increasing gradually to 70 or more, suspending the treatment when intolerance is shown Externally the leprous parts are treated with an ointment of Gurjun Balsam, 3 parts, Lanolin, 1 part—*P J* '95, 11 27

It is used in India as a substitute for Balsam of Copaiba in gonorrhœa, also as a natural varnish

Prescribing Notes—*Best prescribed in capsules for internal administration As a local application in the form of an emulsion made with Lime Water, or as an ointment made with a Lanolin basis*

Descriptive Notes—It is also known as Wood Oil, but must not be confounded with the Wood Oil of China, which is a drying fixed Oil, used in China

instead of Linseed Oil, and is derived from *Aleurites Fordii*, Harms', and other species, it is also known as Tung Oil. To prevent confusion on the name, Gurjun Oil or Gurjun Balsam should be used for Balsamum Dipterocarpi.

BALSAMUM PERUVIANUM.

BALSAM OF PERU

FR, BAUME DU PEROU, GER, PERUBALSAM, ITAL, BALSAMO PERUVIANO, SPAN, BALSAMO DEL PERU

A dark brownish viscid liquid, obtained (by special treatment) from the Trunk of *Miconia Peruvica*, Klotzsch, growing in San Salvador, Central America.

It consists mainly of Cinnamoin, the Benzyl Ester of Benzoic Acid, with a smaller proportion of the Benzyl Ester of Cinnamic Acid, free Cinnamic Acid, traces of Vanillin and Pteroesinotannol Esters of Cinnamic and Benzoic Acid.

Solubility—1 in 1 of Alcohol (90 p.c.), when more than 3 of Alcohol are added to 1 of Balsam it becomes turbid, in all proportions of Chloroform, insoluble in Olive Oil.

Medicinal Properties—Stimulant and disinfectant expectorant. Useful in chronic bronchitis, contra-indicated in acute bronchial catarrh because of its stimulant action, also used as a urinary antiseptic.

Externally as an ointment for chronic indolent ulcers and for sore nipples, for scabies and pediculi and parasitic skin diseases, to relieve itching in urticaria, and prevent or heal bed-sores.

The Balsam contains an Essential Oil, the vapour of which is extremely toxic to the acarus of itch. The patient is rubbed in the evening for fifteen or twenty minutes with the Balsam, it is not necessary to rub hard, as the vapour is sufficient to kill the parasite—*L* '96, i 1101.

As a dressing in warfare, it may be left on for 20 days without removal and disinfection, and sterilisation is unnecessary—*L* '04, ii, 1807.

Not to be applied to large areas of skin for scabies in children, small bodied adults, and patients with renal trouble, as it may produce albuminuria or nephritis—*B M J* '07, i 972.

Superior to sulphur in scabies—*B M J* '07, ii 1710.

Dose.—5 to 15 minims = 0.3 to 0.9 c.c.

Prescribing Notes—Given as an emulsion with Mucilage of Gum Acacia, or Sugar and yolk of Egg with Water.

Not Official.—Unguentum Peruvianum, and Unguentum Peruvianum Resinosum.

Foreign Pharmacopœias—Official in Austr., sp gr 1.14 to 1.16, Belg., 1.187 to 1.150, Dutch, sp gr 1.14 to 1.145, Dan., Ital., Norw. and Swed., sp gr 1.135 to 1.150, Fr., sp gr 1.135 to 1.150, Ger., sp gr 1.140 to 1.150, Hung., sp gr 1.137 to 1.150, Jap., sp gr 1.140 to 1.162, Russ., sp gr 1.135 to 1.145, Port., sp gr 1.15, Span., sp gr 1.13 to 1.16, Swiss, sp gr 1.145 to 1.155, U.S., sp gr 1.140 to 1.150, at 25° C (77° F), Mex., 1.14 to 1.145.

Descriptive Notes—Balsam of Peru is not a natural exudation, but is a pathological product formed after the bark has been beaten and scorched. It is a nearly black, oily liquid, heavier than Water,

and with a balsamic rather smoky odour, which is fragrant and agreeable when the balsam is smeared on paper and warmed. It has not a very pronounced taste, but leaves an unpleasant burning sensation in the fauces. It is liable to be adulterated with several substances, *see* Tests. An artificial Balsam of Peru, called Perugen, is now sold in commerce, but its use in medicine is not justified, since it cannot be said to be derived from *Myroxylon Perense*. With the Nitric Acid test of Cæsar and Loretz, *PJ* (4) xxi, p. 579, Perugen gives an intense olive-green colour, instead of the golden yellow yielded by pure Balsam.

Tests—The distinguishing tests for Peruvian Balsam are (1) the specific gravity, which should be 1.137 to 1.150 [the *USP* gives 1.140 to 1.150 at 25° C (77° F), the *PG* 1.140 to 1.150], (2) the presence of from 57 to 60 p.c. of Cinnamoin as determined by first shaking a weighed quantity of 5 grammes of the Balsam with 5 c.c. of Sodium Hydroxide Solution (15 p.c.) and shaking the latter solution with three successive portions each of 15 c.c. of Ether (sp. gr. 0.720), the residue from the ethereal solution on evaporation being dried until the loss between two weighings at intervals of five minutes does not exceed 0.01 gramme, (3) the percentage of Potassium Hydroxide required to saponify the residue, which should be from 23.23 to 23.76 p.c. as determined by saponifying the weighed residue from the Ether treatment with Normal Volumetric Alcoholic Potassium Hydroxide Solution titrating the uncombined alkali with Normal Volumetric Sulphuric Acid Solution.

The *USP* requires the Balsam to contain at least 56 p.c. of Cinnamoin as determined by a single extraction with Ether from a mixture of the Balsam and Sodium Hydroxide Test Solution, the residue from the Ether treatment should require not less than 23.49 p.c. of Potassium Hydroxide for saponification. The *PG* test indicates 56 p.c. of Cinnamoin as determined by extraction with three successive portions of Ether from a mixture of the Balsam, Water, and Sodium Hydroxide Solution (15 p.c.), the Ether residue should require not less than 23.66 p.c. of Potassium Hydroxide for its saponification. Both *USP* and *PG* employ Semi-normal Volumetric Hydrochloric Acid Solution for titrating the uncombined alkali after saponification. Attention has been called (*PJ* '01, i. 29) to the ambiguous wording of the Pharmacopœia test. The original test directs the exhaustion of the Balsam with Ether and treatment of the filtered ethereal solution with Sodium Hydroxide Solution. The original directions are that the Balsam shall be shaken with Sodium Hydroxide Solution (15 p.c.) and then washed with Ether, the Ether removed and the residue weighed after suitable drying. The residue would be taken to mean the residual Balsam after treatment with Sodium Hydroxide Solution (15 p.c.) and Ether, whereas the residue of Cinnamoin and other Ether soluble bodies is intended.

A determination of the Acid, Ester and Saponification value of the original Balsam is useful in judging of the quality of a sample. The *BP* does not make any reference to such determinations. The *USP*

gives a quantitative test for limit of Acid Resins and according to the test these should not amount to more than 14.09 p.c. reckoned as Cinnamic Acid. No saponification test on the original Balsam is given. The *PG* omits the 'limit of Acid Resins,' but includes a Saponification test, the Balsam being required to neutralise not less than 22.46 p.c. of Potassium Hydroxide, indicating a Saponification value of 224.6.

The more generally occurring impurities are Copaliba Balsam, Colophony, fixed oils, *et c.*, Olive and Castor Oils, Ethylic Alcohol, Turpentine, Storax and Gurjun Balsam. As a general rule adulterants raise the Acid and lower the Saponification values. The *BP* employs trituration with Lime as a test for ensuring the absence of Copaliba and Resins and when warmed until the volatile matter is dissipated and effervescence commences, the absence of fatty odour is assumed to indicate the absence of Castor and other Fatty Oils. No diminution in volume when equal volumes of the Balsam and Water are shaken together indicates the absence of Ethylic Alcohol. The separation of about 40 p.c. of Resin and a clear pale brown supernatant fluid with only a slight fluorescence, when one part of the Balsam is treated with three parts of Carbon Bisulphide, is officially stated to indicate the absence of Gurjun Balsam. The trituration with Lime is included in the *USP*, though the latter part of the test is omitted; the test was official in *PG* III, but is omitted altogether in *PG* IV. Neither *USP* nor *PG* includes a test for Ethyl Alcohol, nor the Carbon Bisulphide test. Both *USP* and *PG* include a Sulphuric Acid test for Fixed Oils and Gurjun Balsam. The *USP* alone includes a test with Benzoin and concentrated Nitric Acid for the detection of Resin, the balance of the Benzoin solution being shaken with Copper Acetate Solution, the non-production of a green or bluish-green coloration indicating the absence of Resin, Turpentine, Storax, Fatty Oils, etc. Dieterich is of opinion that qualitative reactions are devoid of any value in judging the quality of a Balsam, and suggests including a determination of the Resin-ester. The following test has been suggested by Dieterich (*Beichte der deutschen pharmaceutischen Gesellschaft* 18, 135) for inclusion in the *BP* tests.—The alkaline solution of the Resin obtained in the assay of the balsam is acidified and shaken with 10 c.c. of Ether, 5 c.c. of this solution is poured into a test-tube, and after inclining the tube 1 c.c. of concentrated Sulphuric Acid is allowed to flow in, the test-tube is brought into a vertical position, and after a short time is again inclined and 2 c.c. of concentrated Hydrochloric Acid allowed to flow in. A reddish-brown, but not a green or greenish-brown zone should develop between the Hydrochloric Acid and the Ether, a red zone between the Sulphuric Acid and the Hydrochloric Acid. This test is intended to detect the presence of synthetic Balsam of Peru (Perugen).

Calcium Oxide—When the Balsam is mixed with half its volume of Calcium Hydroxide and heating on a water-bath for half an hour a solid mass should not be formed, indicating the absence of Resin, Storax, or Copaliba, *USP*.

Sulphuric Acid—If 10 drops of the Balsam be triturated with 20 drops of Sulphuric Acid a tough homogeneous mass results, which when washed with cold

Water develops a violet colour on its surface, and, after draining out the Water a brittle mass is obtained which may be crumbled up, indicating the absence of fixed oils, *P G* and *U S P*

Benzin and Nitric Acid—If 1 gramme of the Balsam be shaken with 5 c c of Petroleum Benzin, the mixture warmed on a water bath for 10 minutes, and sufficient solvent added to replace loss by evaporation, then if 2 c c of this Benzin Solution be evaporated and treated with a drop of Nitric Acid, *sp gr* 1.42, a permanent green or bluish green colour should not be produced (absence of Resin), *U S P*. The remaining 3 c c of Benzin Solution when shaken with an equal volume of aqueous Solution of Copper Acetate (1-1000) should not be coloured green or bluish green (absence of Resin, Turpentine, Storax, Fatty Oils, etc), *U S P*

Saponification—Let 1 gramme of Balsam be dissolved in 20 c c of Alcohol (90 p c) and 50 c c of Semi-normal Alcoholic Solution of Potassium Hydroxide be added, then let the mixture be heated on a water-bath for half an hour. Dilute with 300 c c of Water and titrate with Semi-normal Solution of Hydrochloric Acid, not more than 42 c c of the Acid solution should be necessary to neutralise the excess of Potassium Hydroxide Solution, *P G*

Volumetric Determination of Free Acid—The *U S P* directs 1 gramme of Balsam to be dissolved in 100 c c of Alcohol (94.9 p c), and titrated with Semi-normal Alcoholic Solution of Potassium Hydroxide, using 1 c c of Phenolphthalein T S as indicator, when not more than 2 c c of the Volumetric Alkali Solution should be required to produce a pink colour

Gravimetric and Volumetric Determination of Cinnamain—The *B P* quantities are outlined in the large type notes above, the *P G* quantities are 2.5 grammes of Balsam, 5 c c Sodium Hydroxide Solution (15 p c w/w) and 5 c c of Water, washing with three quantities of Ether 10 c c each. The final residue should amount to at least 1.4 grammes. The *U S P* directs the mixture of 3 grammes of Balsam with 80 c c Sodium Hydroxide T S, then washing with 60 grammes of Ether, and the careful evaporation of 51.5 grammes of the ethereal liquid, when not less than 1.4 grammes of residue of constant weight should be obtained (presence of at least 56 p c of Cinnamain). The residue obtained from the *B P* gravimetric determination of Cinnamain is dissolved in 40 c c of Alcohol (90 p c), and saponified under a reflux condenser for one hour with 20 c c of Normal Volumetric Alcoholic Potassium Hydroxide Solution, the excess of Volumetric alkali solution being titrated with Normal Volumetric Sulphuric Acid Solution, to neutralise this excess from 7.2 to 8.1 c c should be necessary. The *P G* and *U S P* direct the solution of the weighed residue in 25 c c of Alcohol, mixing with 25 c c Semi-normal Volumetric Alcoholic Solution of Potassium Hydroxide and carefully heating on a water bath for half an hour, then after the addition of 10 drops (1 c c, *U S P*) of Phenolphthalein T S as indicator, the mixture should require not more than 13.2 c c Semi-normal Volumetric Solution of Hydrochloric Acid for exact neutralisation

Not Official

UNGUENTUM PERUVIANUM—Balsam, 1, Lard, 7

This has been incorporated in the *B P C*

UNGUENTUM PERUVIANUM RESINOSUM—Balsam, 1, Resin Ointment, 1

BALSAMUM TOLUTANUM.

BALSAM OF TOLU

FR, BAUME DE TOLU, GFR, TOLUBALSAM, ITAL, BALSAMO TOLUTANO, SPAN., BALSAMO DE TOLU

A yellowish-brown, soft, tenacious mass, which exudes from the Trunk of *Myroxylon Toluifera*, H B and K, on incision. Imported from the northern ports of Colombia, South America

might be inserted in the place of the Potassium Hydroxide in the requirement as to the quantity necessary for saponification. One-third of the weight of the residue is considered to be too high, and it is suggested that the sentence should read not less than 290 parts of Potassium Hydroxide per 1000 parts of dry residue. A saving of time would, moreover, be accomplished by expressing the percentage of Potassium Hydroxide required, on the original Balsam instead of on the residue, the evaporation of the Carbon Bisulphide would then not require to be carried beyond the stage necessary for the removal of the solvent. The Carbon Bisulphide test is not included in the *USP* and *PG*, but both these Pharmacopœias give an Acid and Ester value. According to the *USP* the amount of Potassium Hydroxide required to neutralise the acidity should be not less than 11.14 p.c. nor more than 16.72 p.c. corresponding to an Acid value of 111.4 to 167.2. The *PG* requires not less than 11.23 p.c. nor more than 16.84 p.c. corresponding to an Acid value of 112.3 to 168.4. The amount of Potassium Hydroxide required by the *USP* to saponify the Esters should be not less than 15.32 p.c. nor more than 18.95 p.c. corresponding to a Saponification value of 153.2 to 189.5. That required by the *PG* is not less than 15.44 p.c. nor more than 19.09 p.c., corresponding to a Saponification value of 154.4 to 190.9.

The more generally occurring impurities are Resin and Copaiba. The *BP* relies solely on the sufficient proportion of Benzoates and Cinnamates as indicated by the Carbon Bisulphide residue and its Saponification figure, the *PG* on the Acid and Saponification value, whilst the *USP* gives in addition confirmatory qualitative tests with Sulphuric Acid on a solution in Glacial Acetic Acid of residue obtained on evaporating a Carbon Bisulphide solution of the Balsam, and the non-production of a green coloration when a 1 in 8 solution of the Balsam in Benzine is shaken with an equal volume of a 1 in 1000 aqueous Copper Acetate solution.

The following test has been suggested by Dieterich for the detection of Colophony.—A weighed quantity of 0.5 gramme of the balsam is mixed with 5 c.c. of Water, and 5 c.c. of Sodium Hydroxide Solution (15 p.c. w/w), the mixture is shaken with 10 c.c. of Ether, separated, acidified, and again shaken with Ether. A measured portion of the ethereal solution is introduced into a test-tube, the tube inclined and 1 c.c. of concentrated Sulphuric Acid allowed to flow in, again brought into a vertical position, and after a short time again inclined, and 2 c.c. of Hydrochloric Acid allowed to flow in. It should produce a red zone between the Hydrochloric Acid and the Ether, and a deep red zone between the Sulphuric Acid and the Hydrochloric Acid. A weighed quantity of 1 gramme of the balsam dissolved in 5 c.c. of Glacial Acetic Acid and 2 drops of Sulphuric Acid dropped into the solution, previously heated to boiling, should produce a bluish-violet or bluish-green colour.

Carbon Bisulphide, Glacial Acetic, and Sulphuric Acids.—If 0.5 gramme of the Balsam be shaken with 25 c.c. of Carbon Bisulphide, allowed to stand for 30 minutes, filtered, and the filtrate evaporated to dryness, a residue is

obtained which when dissolved in Glacial Acetic Acid should not yield a green colour on the addition of a few drops of Sulphuric Acid, *U S P*

Benzin and Copper Acetate—If 1 gramme of the Balsam be agitated with 8 c.c. Petroleum Benzin for 5 minutes, the supernatant liquid should not be coloured green when shaken with an equal volume of a 1 in 1000 solution of Copper Acetate, indicating the absence of Resin and Copaiba, *U S P*

Acid Value—If 1 gramme of the Balsam be dissolved in 50 c.c. of Alcohol and 10 drops (1 c.c. *U S P*) of Potassium Permanganate Solution be added, then the addition of not less than 4 c.c. and not more than 6 c.c. of Semi-normal Volumetric Alcoholic Solution of Potassium Hydroxide should be required to produce a red colour, indicating the limit of acidity, *P G* and *U S P*

Saponification Value—If this liquid be mixed with more Semi-normal Alcoholic Solution of Potassium Hydroxide until the total amount used is 20 c.c., and the mixture heated on a water-bath for half an hour and allowed to cool, then on titrating with Semi-normal Volumetric Solution of Hydrochloric Acid (Semi-normal Volumetric Solution of Sulphuric Acid, *U S P*) the liquid should require not less than 13.2 and not more than 14.5 c.c. of the Acid Solution to neutralise the excess of Potassium Hydroxide Solution, *P G* and *U S P*.

Preparations.

SYRUPUS TOLUTANUS. SYRUP OF BALSAM OF TOLU

FR., SIROP DE BAUME DE TOLU, GER., TOLUBALSAMSIROP, ITAL., SCIROPO DI BALSAMO DEL TOLU, SPAN., JARABE DE BALSAMO DE TOLU

Balsam of Tolu, 1½, is boiled with 20 of Distilled Water to produce 16 of liquid, in which (after filtration) are dissolved 32 of Sugar. When finished it should weigh 48

A better flavoured Syrup may be made as follows—Balsam of Tolu, 1½, Sugar, 8 Powder the Tolu with the Sugar, macerate in Water 16, for 24 hours, with frequent agitation, filter bright, and dissolve in it (cold) Sugar 24

In a paper read before the British Pharmaceutical Conference (*P J*, '99, ii 103, 116, *C D* '99, ii 212, 228) the superiority of a Syrup made on the above lines has been clearly demonstrated. Six Syrups were prepared, the two samples of genuine Balsam used being previously submitted to careful examination. The Syrup prepared by the *Companion* process ranked first, having the best flavour and containing 1.12 parts of Cinnamic Acid out of 3.33 contained in the Balsam used. The *B P* Syrup ranked last, containing in summer 1.00, and in winter 0.60 parts of Cinnamic Acid

Syrupus Tolutanus (*U S*)—Tincture of Tolu, 5, is triturated with Magnesium Carbonate, 1, and Sugar, 6, gradually adding Water 45, filtering, dissolving Sugar 76 in the liquor with aid of gentle heat, straining while hot and adding sufficient Water to make 100

Dose—½ to 1 fl. drm. = 1.8 to 3.6 c.c.

Liquor Tolutanus pro Syrup.—Balsam of Tolu 4, dissolved in 12 of Alcohol (90 p.c.) and added to 26 of Water previously heated to 70° C. and placed in a bottle, shaken vigorously, then set aside for 24 hours, filter bright. To make Syrup of Tolu mix 1 of the solution with 7 times its volume of simple syrup—(*Farr and Wright*) *Y B P* 1899, 366

This has been incorporated in the *B P C*.

Foreign Pharmacopœias—Official in Belg., Fr., Ital., Norw., Port., Span.; Swed., Swiss and U.S., Mex. made with Tincture. Not in the others

Belg. and Fr. have Tablets of Balsam of Tolu.

TINCTURA TOLUTANA. TINCTURE OF BALSAM OF TOLU.

1 of Balsam of Tolu, macerated with Alcohol (90 p.c.) 95 to yield 10.
(1 in 10)

Dose — $\frac{1}{2}$ to 1 fl drm = 1·8 to 3·6 c c

Foreign Pharmacopœias —Official in Belg, Dan, Fr, Swed and U S,
1 in 5, Port, 8 in 20 All by weight, except U S Not in the others

Tests —Tincture of Tolu has a sp gr of 0·860 to 0·865, it contains from 8 to 10 p c w/v of total solids and about 80 p c of Absolute Alcohol by volume

Not Official

BAPTISIN

A powdered extract obtained from *Baptisia tinctoria*, R. Di In small doses, laxative, in large doses, purgative and emetic

Dose —1 to 5 grains = 0·06 to 0·32 gramme Usually given in pill

Not Official

BARI SULPHIDUM

It is somewhat difficult to obtain in a pure condition, and commercial samples as a rule do not contain more than 50 p c BaS

Medicinal Properties —The chief use of this is as a depilatory, for which purpose it is unequalled, removing hair with less injury to the skin than any other application

Test —For the estimation of BaS 1 Make a standard Zinc Solution by dissolving 7·7 grammes of Zinc in about 75 c c of Diluted Hydrochloric Acid, adding excess of Ammonia Solution and diluting to 1000 c c, 2 Make an alkaline Lead Solution by dissolving 1 gramme Lead Acetate in about 20 c c of hot Solution of Potassium Hydroxide and diluting to 100 c c, 3 Heat to boiling 1 gramme of Barium Sulphide in about 50 c c of Water and titrate with the standard Zinc Solution till no black or brown colour is obtained by adding a drop of the filtered Barium Solution to a drop of the Lead indicator, spotted on a porcelain slab Each c c of the Zinc Solution used is equivalent to 2 per cent of Barium Sulphide in the sample operated upon

Preparation

DEPILATORY —Barium Sulphide (containing 70 p c BaS, or an equivalent quantity of any other strength), in fine powder, 2, Starch, 5, Orris Root, in powder, 1 Mix

For use make it into a thin paste with Water, apply to the part from which the hair is to be removed, after five minutes scrape off with a blunt knife

Not Official

BARI CHLORIDUM

BaCl₂, eq, 206·78

Colourless crystalline plates

Solubility —1 in 2½ of Water

Medicinal Properties —Occasionally given in syphilis, scrofula, and as a cardiac tonic, but requires care on account of its toxic properties

Dose — $\frac{1}{2}$ to 2 grains = 0·016 to 0·13 gramme

Toxic effects occur only with large doses —J. '03, i 1194

Stated (B M J E. '05, i 24) to have no cumulative effect in ordinary doses, does not disturb the digestion or irritate the kidneys Usual adult dose, 3 grains = 0·2 gramme, 3 or 4 times daily in Water.

Foreign Pharmacopœias—Official in Ger, Mex. and Swiss. Not in the others

Tests—An aqueous solution yields with diluted Sulphuric Acid Solution or an aqueous solution of a soluble Sulphate, a heavy white precipitate insoluble in concentrated Hydrochloric Acid and in strong Nitric Acid, with Potassium Bichromate Solution it yields a yellow precipitate soluble in diluted mineral acid. With Silver Nitrate Solution it yields a white curdy precipitate insoluble in Nitric Acid, soluble in Ammonia Solution, and in Potassium Cyanide Solution. It should yield no coloration or precipitate when acidified with Hydrochloric Acid and tested with Hydrogen Sulphide Solution. The P.G. includes a test for Iron, 20 c.c. of a 1 in 20 aqueous solution shall not be coloured blue by 0.5 c.c. of Potassium Ferrocyanide Solution (5 p.c.) No residue shall remain after complete evaporation of the Barium by Sulphuric Acid, evaporation of the filtrate and ignition at a low red heat

Not Official

BEBEERINÆ SULPHAS

Dark-brown thin translucent scales, yellow when in powder, with a strong bitter taste. A preparation made from *Nectandra* or *Bebeeru* Bark (*Nectandra Roduei*, Schomb.), containing about 60 p.c. of alkaloids, one half being **Bebeerine** (Beberine), $C_{17}H_{21}NO_3$, eq. 308.87. It was official from 1861 till 1898.

Solubility—Sparingly in Alcohol (90 p.c.), dissolves about 1 in 1 of Water, and the solution can be diluted up to 1 and 8 of Water, but on further dilution it precipitates until about 80 or 100 parts of Water have been added, but samples vary in this respect, readily soluble in Water containing a mineral Acid.

Medicinal Properties—Aromatic bitter, stomachic tonic, an imperfect substitute for Quinine.

Dose.—1 to 5 grains = 0.06 to 0.32 gramme.

Prescribing Notes—*Given in solution, or in pills made with 'Dispensing Syrup.'*

The following pure products are commercial. **Bebeerine** pure, slightly soluble in Water, readily in Alcohol, Chloroform and Ether, **Bebeerine Hydrochloride** and **Bebeerine Sulphate**, are both readily soluble in Water and Alcohol. Dose of the two latter 1 to 2 grains = 0.06 to 0.13 gramme.

Not Official

BELÆ FRUCTUS.

Bael Fruit is obtained from *Ægle Marmelos*, Correa.

The dried half-ripe Fruit was formerly official, but is now omitted.

The fresh half-ripe Fruit is now official in the *Ind* and *Col. Add.* for use in India and the Eastern Colonies, as is also the Liquid Extract.

Medicinal Properties—The Fresh Fruit has been much extolled in India for diarrhoea and dysentery, and the Confection prepared in Britain appears to have similar properties. The Dried Fruit is not considered a trustworthy remedy.

If the fresh fruit is not obtainable, the official preparation **Extract Belæ** Liquid answers well in the treatment of dysentery. Generally speaking, Milk is the best vehicle for Bael Fruit and pulp.—*I.M.G.* '05, ii 264.

CONFECTIO BELÆ RECENTIS (*Synure*)—Prepared from Fresh Fruit, imported from India in the Spring months. It retains the odour and flavour of the Fresh Fruit.

Dose—A teaspoonful.

EXTRACTUM BELÆ LIQUIDUM (*Ind* and *Col. Add.*)—Made by macerating 4 of bruised Bael Fruit in Water by successive treatments, evaporating the mixed fluids to 3, and when cold adding Alcohol (90 p.c.), *q.s.* to make 4.

Dose—1 to 2 fl. drms = 3.6 to 7.2 c.c.

BELLADONNA.

BELLADONNA

FR, BELLADONE, GER, BELLADONNA, ITAL, BELLADONNA, SPAN, BELLADONA

The fresh Leaves and Branches of *Atropa Belladonna*, L, as well as the dried Root, are official

Medicinal Properties—Anodyne, antispasmodic, mydriatic, antilactagogue, anhydrotic, and urinary sedative. There is no drug which can compare with it in checking the secretions of milk, sweat, and saliva. It is given for epilepsy, and is one of the best remedies for whooping-cough and for painful spasm of the bladder, in renal colic, dysmenorrhœa and typhlitis, in full and frequent doses for asthma, both as a prophylactic and curative. It is of the utmost value in relieving cardiac pain and distress, palpitation and aortic regurgitation. Useful in typhoid with contracted pupil, and in acute bronchitis it stops profuse secretion. In large or continued doses it causes dilatation of the pupil and dryness of the mouth and throat. For habitual constipation $\frac{1}{4}$ to $\frac{1}{2}$ grain of Extract on rising in the morning. For nocturnal incontinence of urine, 5 to 10 minims of the Tincture, with the same dose of Tinct of Perchloride of Iron three times a day. Ringer recommends larger doses of Belladonna for this troublesome complaint in children, 10 to 30 minims of the Tincture three times a day, small doses often fail when large doses at once succeed. Useful in loss of tone and irritable state of the generative organs which gives rise to nocturnal emissions, although it has slightly aphrodisiacal properties.

For external uses, see Prescribing Notes.

In the treatment of epilepsy (L '05, i 710) it is a remedy which should be tried in all cases in which the Bromides have failed. Every now and again cases will be met with in which this drug produces remarkable and persistent arrest of seizures.

Cases of poisoning (L '05, i 714) by application of Belladonna plaster 8 x 5 applied to the loins for the relief of lumbago—Recovery. Ointment of 4 grains Atropine Sulphate in the oz applied for chronic eczema of the nates causes toxic symptoms—Recovery.

Poisoning by liniment, recovery after use of Strychnine—B M J '07, ii 1515.

Prescribing Notes—The Extract in pills, also the Tincture and Succus are for internal use. The Suppository is used in prostatitis, cystitis and chordee. Bougies made with Gelatin base or Oil of Thiodon contain $\frac{1}{4}$ to 2 grains of Alcoholic Extract of Belladonna. Externally the Liniment and Compound Liniment sprinkled on impermeable Plumb are very useful in pleurodynia, lumbago and muscular rheumatism, as is also the Chloroform preparation alone or mixed with oil. The Glycerinum as a paint, and the Emplastrum, are used for sprains, acute synovitis, and to check mammary secretion and prevent inflammation of the breast, both are also an excellent remedy in cardiac pain and palpitation. Extract of Belladonna is a component of many Hospital formulas for pills, and is prescribed with Aloes, Camphor, Quinine, Rhubarb, Valerian and Zinc Oxide, in doses of $\frac{1}{2}$ to $\frac{1}{4}$ grain in each pill. In Eye Lotions 2 grains of the green extract to the fl oz.

Dose.—Will be found under the respective preparations.

Incompatibles.—Caustic Alkalis, Opium, Strychnine.

Official Preparations—*Extractum Belladonnæ Vinide*, and *Succus Belladonnæ*, from the fresh leaves and branches *Extractum Belladonnæ Liquidum*, from the dried root. *Emplastrum Belladonnæ*, *Extractum Belladonnæ Alcoholicum*, *Linimentum Belladonnæ*, *Tinctura Belladonnæ*, and *Unguentum Belladonnæ*, from the **Liquid Extract**, *Suppositoria Belladonnæ*, from the **Alcoholic Extract**. *Atropine*, from leaves or root

Not Official—*Chloroformum Belladonnæ*, *Collodium Belladonnæ*, *Emplastrum Belladonnæ Vinide*, *Extractum Belladonnæ Foliorum*, *Extractum Belladonnæ Exsiccatum*, *Glycerinum Belladonnæ*, *Linimentum Belladonnæ Compositum*, and *Ethereal Tincture of Belladonnæ*

Antidotes.—In cases of poisoning by Belladonna, use stomach-tube or give one of the following emetics: 10 grains of Copper Sulphate, 20 grains of Zinc Sulphate, 1 oz of Ipecacuanha Wine, or hypodermic injection of $\frac{1}{10}$ grain Apomorphine. Give stimulants, inject Pilocarpine, an enema of Coffee. If necessary apply artificial respiration.

BELLADONNÆ FOLIA. BELLADONNA LEAVES.

FR, FEUILLES DE BELLADONNE, GER, BELLADONNABLAUFLER, ITAL, FOGLIE DI BELLADONNA, SPAN, HOJA DE BELLADONA

The fresh Leaves and Branches of *Atropa Belladonna*, L., collected when the plant is in flower

The Leaves are official in the *USP* and *PG*. The *USP* has introduced a process of assay and requires that they shall yield not less than 0.33 p.c. of mydriatic alkaloids when assayed by this process. Neither *BP* nor *PG* fixes a standard for the Leaves

Descriptive Notes—The Pharmacopœia directs that the fresh Leaves and Branches of *Atropa Belladonna* should be collected when the plant is in flower. But the plant often continues to flower long after the fruits are ripe, and consequently the leaves are likely to vary in strength, it would, perhaps, have been better to use the words 'commencing to flower.' The flowering branches are easily recognised by the dull purple bell-shaped flowers, and the ovate entire leaves apparently arranged in pairs, of which one leaf is smaller than the other. The smaller leaf is, however, a bract belonging to the flower, which is placed outside, not in, the axil of the larger leaf. The lower stem leaves are alternate and not in pairs. The leaves vary in size from 3 to 8 inches (7.5 to 20 cm.) in length, and 2 to 3½ inches (5 to 9 cm.) broad and are glabrous or nearly so.

It will be noted that the dried leaves are not official in the *BP* although they are in the *USP* and the *PG*. The dried leaves are usually brownish-green above and paler beneath, and present, especially on the under surface, when seen under a good lens, minute pale dots or prominences caused by cells filled with sandy crystals of Calcium Oxalate, which do not contract in drying. These crystal cells are easily seen under the microscope in a fragment of the leaf cleared by Chloral Hydrate Solution, as well as the striations of the epidermal cells, which are also characteristic. The dried leaves of *Scopolia carniolica*, Jacq., have been offered in commerce when Belladonna leaves were scarce and dear, but they are thinner, darker green, and the small veins are more prominent. The leaves of *Phytolacca decandra*, L., from Bosnia, have also been offered as Belladonna in European commerce. Their upper surface has no hairs, and is of a

lighter green, and contains no crystal sand, but acicular raphides, and the epidermal cells are polygonal, not sinuate as in *Belladonna*.

The *P G* leaves are collected in the flowering season.

The percentage of alkaloid varies considerably, a good well dried sample should contain about 0.5 p.c.

Ph. Ger. maximum single dose, 0.2 gramme, maximum daily dose, 0.6 gramme.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Et, Ger, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U.S., Leaves, Port, Herb. Not in Hung.

The *Brussels Conference* (1906) uses only the leaf, dried, powdered drug to be used entire.

Tests—No standard for *Belladonna Leaves* is given in either *B.P.* or *P.G.*, the *U.S.P.* states that they should yield not less than 0.33 p.c. of mydriatic alkaloids when assayed by the *U.S.P.* process. The *U.S.P.* method of standardisation is as follows—A weighed quantity of 10 grammes of the leaves in No. 60 powder is allowed to stand for ten minutes in an Erlenmeyer flask with 50 c.c. of a mixture containing 4 parts by volume of Ether and 1 part by volume of Chloroform. A mixture of 2 c.c. of Ammonia Water with 3 c.c. of Water is added and the whole shaken at frequent intervals for one hour. The contents of the flask are then transferred, as far as possible, to a small percolator inserted in a separator containing 6 c.c. of Normal Volumetric Sulphuric Acid Solution diluted with 20 c.c. of Water. After the liquid has passed through, the leaves are packed in the separator, the flask washed first with 10 c.c. and then with several portions of 5 c.c. of the Chloroform-Ether mixture, and these with the remaining contents of the flask transferred to the percolator, the percolation being continued with the Chloroform Ether mixture until 50 c.c. has been used. The separator is now securely stoppered, agitated for one minute, the fluids allowed to separate clear, and the acid aqueous solution removed to a second separator. A further quantity of 10 c.c. of a mixture of Normal Volumetric Sulphuric Acid Solution of the same strength as that previously used is added, the contents well shaken, allowed to separate and the acid aqueous portion again drawn off into the second separator, and this operation is repeated. To the mixed acid liquids in the second separator is added a small piece of red Litmus paper and sufficient Ammonia Water to render the fluid distinctly alkaline, the mixture being then shaken with two successive quantities of 15 c.c. and one of 5 c.c. of Chloroform. The Chloroform solutions are separated, collected, and evaporated in a beaker, the residue dissolved in 3 c.c. of Ether and the Ether allowed to evaporate. The residue is dissolved in 3 c.c. of Tenth-normal Volumetric Sulphuric Acid Solution, 5 drops of Cochineal Test Solution added and the excess of Acid titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution. The number of c.c. of Fiftieth-normal Volumetric Potassium Hydroxide Solution required is divided by 5, the quotient subtracted from 3 and the product multiplied by 0.0287 and again multiplied by 10, giving the percentage of total mydriatic alkaloids in the leaves.

EXTRACTUM BELLADONNÆ VIRIDE. GREEN EXTRACT OF BELLADONNA.

Heat the expressed Juice of the Fresh Leaves and Young Branches to 54·4° C (130° F), and collect on a filter the green colouring matter which separates, heat the filtrate to 93·3° C (200° F) and filter. Evaporate the clear liquid to an Extract, returning the green colouring matter towards the end of the process and completing the operation at 60° C (140° F)

100 lb of Herb yielded 56 lb of Juice, or nearly 4 lb Extract

100 lb Leaves, when dried, weighed 16 lb

An estimation of the alkaloids contained in four samples of Extract of Belladonna, prepared in 1885 by different makers, gave 0·94 p.c., 1·17 p.c., 1·11 p.c., and 0·73 p.c. The following samples in good condition were examined at the same time 1880—1·26 p.c., 1·22 p.c., 1881—1·16 p.c., 1·21 p.c., 1884—1·21 p.c.

A sample of 1892 Extract yielded 1·7 p.c. of alkaloids

An average sample of Extract contains rather over 1 p.c. of alkaloids

Dose.— $\frac{1}{4}$ to 1 grain = 0·016 to 0·06 gramme.

Ph. Ger. maximum single dose, 0·05 gramme, maximum daily dose, 0·15 gramme

Foreign Pharmacopœias—Official in Austr., Belg. and Mex., Alcoholic from the leaves, Mex. has also Alcoholic extract from the root, and a Fluid Extract 1 in 1. Dan., made from leaves with weak Spirit, Dutch, Alcoholic from leaves, Fr., Alcoholic from leaves and seeds, Ger., made with Water and Spirit from leaves, Hung. and Ital., Alcoholic from root, Norw. and Swed. Alcoholic from leaves, Port., aqueous from dried leaves, Alcoholic from fresh herb and Alcoholic extract purified by Alcohol, Russ., made from leaves with Water and Spirit, Span., aqueous from fresh leaves, also Alcoholic from dried leaves, U.S.—an Alcoholic extract from the powder of the leaf, also Fluid Extract of the root

Extractum Belladonnæ Foliorum (U.S.)—To be prepared with a mixture of Alcohol (95 p.c.) 2, and Water 1, and adjusted with powdered Sugar of Milk to contain 1·4

Extractum Belladonnæ Exsiccatum—Syn. **Extractum Belladonnæ Folii Exsiccatum**. Belladonna Leaves (as per test with Alcohol (70 p.c.) and adjusted with Powdered Belladonna Leaves so as to finally form a powder containing 1 p.c. of alkaloids—*B.P.C.*

The *Brussels Conference* (1906) prepares a solid extract (containing about 10 p.c. of Water) by means of Alcohol (70 p.c.)

The Belg., Dan., Fr. and Swiss Ph. adopt this standard

Tests.—Green Extract of Belladonna, *B.P.*, is not a standardised preparation. The official alcoholic Extract is made from the standardised Fluid Extract, and is required to contain 1 p.c. of the alkaloids of Belladonna Root. The method of procedure adopted by the *U.S.P.* for the standardisation of the Extract from the leaves is as follows—A weighed quantity of 5 grammes of the Extract is dissolved in a mixture of 5 c.c. of Alcohol (94·9 p.c.), 10 c.c. of Water, 2 c.c. of Ammonia Water, and 20 c.c. of Chloroform, and transferred to a separator, using a little Alcohol (94·9 p.c.) to wash out the vessel in which the Extract was dissolved. After the contents of the separator have been well shaken for half a minute they are allowed to separate, the chloroformic solution is removed to another separator, and the contents of the first separator are shaken with another 10 c.c. of Chloroform. After a similar period of shaking and allowing the

liquids to separate, the chloroformic solution is transferred to the second separator. This process is repeated with a further quantity of 10 c c of Chloroform. The alkaloids are then extracted from the mixed chloroformic solutions by shaking for half a minute with 5 c c of Normal Volumetric Sulphuric Acid Solution and 10 c c of Water. The chloroformic layer is removed and again shaken for half a minute with 1 c c of Normal Volumetric Sulphuric Acid Solution and 10 c c of Water, the liquids allowed to separate, and the chloroformic layer removed and rejected. The mixed acid aqueous solutions are filtered through a plug of Cotton wool, the vessels in which they were contained being washed with about 10 c c of Water. After the addition of 15 c c of Chloroform and sufficient Ammonia Water to produce a distinctly alkaline reaction, the contents of the separator are shaken for half a minute, allowed to separate, and the chloroformic layer is drawn off into a beaker. The shakings are repeated, using two separate portions each of 10 c c of Chloroform, the mixed chloroformic liquids are evaporated to dryness, the residue is dissolved in 3 c c of Ether and again evaporated. It is now dissolved in 5 c c of Deci-normal Volumetric Sulphuric Acid Solution, 5 drops of Cocineal or Iodeosin Test-solution added, and the excess of Sulphuric Acid is titrated back with Fiftieth-normal Volumetric Potassium Hydroxide Solution. The number of c c of Fiftieth-normal Volumetric Potassium Hydroxide Solution used is divided by 5, and the quotient is subtracted from 5, the remainder is multiplied first by 0.0287 and then by 20, which gives the percentage of mydriatic alkaloids.

The process of the *P G* is as follows —A weighed quantity of 2 grammes of the Extract is dissolved in 5 grammes of Water and 5 grammes of Absolute Alcohol. 50 grammes of Ether and 20 grammes of Chloroform are added to the solution, and after brisk agitation 10 c c of a 1 in 3 Sodium Carbonate Solution. After the mixture has been allowed to stand for one hour, with frequent intervals of brisk shaking, 50 c c of the clear Chloroform-Ether solution is filtered through a dry, well-covered filter into a flask, and one-half distilled. The remaining Chloroform-Ether solution is transferred to a separator, the flask being washed out with three separate quantities each of 5 c c Ether, and the mixed fluids thoroughly shaken with 20 c c of Hundredth-normal Volumetric Hydrochloric Acid Solution. When the liquids have completely cleared, and after the addition of sufficient Ether to cause the Chloroform-Ether solution to float on the surface of the acid fluid, the latter is filtered through a small filter paper moistened with Water into a flask of white glass holding about 200 c c. The Chloroform-Ether solution is shaken with three successive quantities each of 10 c c of Water, these being passed through the same filter, the filter washed with Water, and the filtrate and washings diluted to about 100 c c. Sufficient Ether to form a layer of 1 cm is added, 5 drops of a 1 in 500 alcoholic Iodeosin solution, and Hundredth-normal Volumetric Potassium Hydroxide Solution until the lower aqueous layer acquires a pale reddish coloration, the liquids being vigorously shaken after each addition. Not more than 13.0 c c of Hundredth normal Volumetric Potassium

Hydroxide Solution should be required. The number of c.c. of Hundredth-normal Volumetric Potassium Hydroxide Solution required, subtracted from 20, multiplied first by 0.0028927, then by 100, and divided by 1.333, will give the percentage by weight of mydriatic alkaloids present in the Extract, which should amount to not less than 1.5 p.c.

The quantity chosen for the assay is a somewhat inconvenient one, the instructions given in the process necessitating the employment of 1.333 as a factor in calculating the percentage yield of alkaloid, the point has already been referred to in Meick's Annual Report for the year 1900. A stock of Chloroform-Ether mixture should be employed instead of a freshly-prepared mixture of Chloroform and Ether, an appreciable rise of temperature ensuing when the liquids are mixed in the indicated proportions. Great attention to the cleanliness of all the vessels employed is also necessary in dealing with Hundredth-normal Volumetric Solutions.

SUCCUS BELLADONNÆ. JUICE OF BELLADONNA

Add 1 of Alcohol (90 p.c.) to 3 of the expressed Juice from the Fresh Leaves and Young Branches.

Dose.—5 to 15 minims = 0.3 to 0.9 c.c.

Belladonna Juice, which would yield an Extract containing 1 p.c. of alkaloid, would form a Succus of about 0.05 p.c., which is also the strength of the Tincture.

BELLADONNÆ RADIX. BELLADONNA ROOT

FR., RACINE DE BELLADONNE, GER., BELLADONNAWURZEL, ITAL., RADICE DI BELLADONNA, SPAN., RAIZ DE BELLADONA

The Root of *Atropa Belladonna*, collected in the autumn and dried.

A good parcel of Roots should average 0.5 p.c. of alkaloid, but occasional bales are found averaging 0.7 to 0.8 p.c.

Foreign Pharmacopœias. Official in Austr., Hung., Ital., Mex., Port., Span., Swiss, Belg., Dan., Dutch, Fr., Ger., Jap., Norw., Russ. or Swed.

Descriptive Notes.—The Belladonna root of commerce is either of English or European origin. That prepared in this country is usually derived from the cultivated plant, the roots of which are dug up every third and fourth year after having furnished Belladonna leaves during that period. It varies in quality, some samples consisting largely of the woody rootstock or underground stem crowning the roots, distinguished by its radiate woody zone, and by being more or less hollow, others consist almost entirely of the true root with occasionally pieces of the horizontal underground stems or suckers which show traces of leaf scars and buds at intervals. The true root, which is the official part, is cylindrical, it may vary considerably in size, but those of medium thickness are to be preferred. It is stated officially to be $\frac{1}{2}$ to $\frac{3}{4}$ in (9 to 19 mm.) in diameter and 6 to 12 in (15 to 30 cm.) or more in length. Vogl gives it as about 10 cm. long, and 1 to 2 cm. thick. Hanbury and Flückiger recommend roots which

are not thicker than the finger. It is of a pale brownish colour with occasional short transverse scars and is finely wrinkled longitudinally, and the surface is easily abraded with the finger nail showing the white starchy tissues beneath. The central portion, clearly defined by the darker cambium ring, is not radiate in structure, but the vascular bundles and vessels are small and scattered irregularly, although more numerous near the cambium. Under the microscope the scattered parenchymatous cells, filled with sandy crystals of Calcium Oxalate, and the starch grains usually in groups of 2 to 3, but rounded when free, form distinctive characters. The root is sometimes met with of darker colour internally and harder, due to slight charring during drying. From Germany it is sometimes imported in carefully dried thin longitudinal slices, so that the official characteristics are not easily seen, and dependence must then be placed on the presence of the sandy raphides, and the form and size of the starch granules. The *U S P* test of a 0.5 percentage of mydriatic alkaloids is not given in the *B P* but would be useful, especially for the sliced root.

The 'root' of *Scopolia Japonica*, Max, sometimes offered as Japanese Belladonna and that of *S. Carniolica*, Jacq, are both characterised by being prostrate rhizomes with numerous closely placed circular stem scars on the upper surface. The true roots, of which there are usually few present, are smaller than Belladonna root and only 2 to 4 inches (5 to 10 cm) long and tapering, and the epidermis is not easily abraded by the nail. Recently Belladonna root imported from Austria has been found to be largely adulterated with the root of *Phytolacca decandra*, L, which though similar in colour is easily recognised by the concentric rings of woody tissue. It is a dangerous adulterant, as it is irritant, acrid and emetic.

Tests—The *B P* does not include a process for the assay of Belladonna Root. The *U S P* gives a process for the determination and requires that it shall yield not less than 0.45 p.c. of mydriatic alkaloids, with the exception that 10 grammes of the root in No. 60 powder are used in the place of 10 grammes of the Leaves, the process is essentially that described under Belladonna Leaves.

The root is not official in the *P G*, but the greater part of the root used in this country comes from Germany.

EMPLASTRUM BELLADONNÆ BELLADONNA PLASTER

4 of Liquid Extract of Belladonna evaporated to 1 and mixed with 5 of Resin Plaster

Contains 0.5 p.c. of alkaloids

This Plaster can be obtained spread on calico, linen, or leather, it is also supplied in rubber combination, spread on felt or kid, plain or porous.

Foreign Pharmacopœias—Official in *F 1*, Extract 1 in 4, *Max*, Belladonna Leaves 20, Alcohol (90 p.c.) 10, *Ammonia* 1, Yellow Wax 80, Turpentine 25, Colophony 25, *Port*, Alcoholic Extract 1, Lead Plaster 9, *Span*, Extract about 1 in 5, *Swiss*, Belladonna extract 1, *Elemi* 1, Colophony 2, Adhesive Plaster 6, *U S*, Extract of Leaves 1, Adhesive Plaster 2½, *Mex*, from the leaves with Alcohol. Not in the others.

EMPLASTRUM BELLADONNÆ (U S)—Standardised Alcoholic Extract of Belladonna Leaves 3, Adhesive Plaster 7. Contains not less than 0.38 p.c. nor more than 0.42 p.c. of mydriatic alkaloids.

EMPLASTRUM BELLADONNÆ VIRIDE (B P C)—Green Extract of Belladonna 1, treated with 4 of Alcohol (90 p.c.), evaporation of the Alcohol, and admixture of the residue with Resin Plaster *q s* to make 4.

It is not standardised and even when made from herb of good quality will only be about half the strength of the official plaster.

Tests.—The Emplastrum Belladonnæ of the *B P* is prepared with the official standardised Fluid Extract of Belladonna, and is officially stated to contain 0.5 p.c. of the alkaloids of the root. The figure is arrived at by calculation, but no process is given by which the presence of the requisite amount of alkaloids may be ensured. One or two processes have been recommended (*P J* '99, ii 110, 114, 180, *C D* '99, ii 214, 227, 331). One of the best and most generally employed is that suggested by Bnd (*P J* '99, ii 146, *Analyst* '99, 175), which is based upon the disintegration of the plaster by Chloroform and Acetic Acid, the removal of the Lead as an insoluble Sulphate, the extraction of the impure alkaloids from the filtered aqueous acid liquid by treatment with Ammonia Solution and Chloroform and then final purification as in the *B P* process for Extractum Belladonnæ Liquidum.

The details of the process are as follows.—A weighed quantity of 15 grammes of the Plaster is gently warmed with 35 c.c. of Chloroform and 5 c.c. of Glacial Acetic Acid until dissolved, and after the addition of a mixture of 35 c.c. of a 1 in 12 dilute Sulphuric Acid Solution and 40 c.c. of Water is again gently warmed and filtered under pressure through a Buchner's filter. The cake of insoluble Lead Sulphate is disintegrated, warmed with a mixture of 10 c.c. of Chloroform, 5 c.c. of dilute Sulphuric Acid, and 10 c.c. of Water, and again filtered. The mixed filtrates are transferred to a separator, the chloroformic layer is removed and washed with two successive quantities of a mixture of 1 c.c. of dilute Sulphuric Acid and 4 c.c. of hot Water, the washings being returned to the aqueous portion. A measured quantity of 20 c.c. of Chloroform and a decided excess of Ammonia Solution are added to the mixed acid aqueous solutions, the mixture gently warmed and agitated. The chloroformic solution is drawn off, and to ensure the complete extraction of the alkaloids the agitation is twice repeated with 10 c.c. of Chloroform. The chloroformic solutions are in each instance separated, mixed and shaken with 5 c.c. of a mixture of 1 in 12 dilute Sulphuric Acid Solution and twice its volume of warm Water. When separation is complete the chloroformic layer is removed and again treated with 5 c.c. of the same mixture as above, the chloroformic layer being again removed, the acid aqueous liquids mixed together and washed with Chloroform, using about 3 c.c. Sufficient Ammonia Solution is then added to form a distinct excess, and the liberated alkaloids are then removed from the mixture by shaking three times in succession with 10 c.c. of Chloroform, the mixed chloroformic solutions, after washing with 5 c.c. of Water containing one drop of Ammonia Solution, are evaporated to dryness on a water-bath in a tared basin, and the

residue dried at a temperature below 100°C (212°F) The amount of alkaloid in this residue may then be determined volumetrically by dissolving in 10 c c of Deci-normal Volumetric Hydrochloric Acid Solution and titrating the excess of acid with Centi-normal Volumetric Sodium Hydroxide Solution, using Tincture of Cochineal, Methyl Orange, Hæmatoxylin, or Iodeosin Solution as an indicator of neutrality The number of c c of Centi-normal Volumetric Sodium Hydroxide Solution used, deducted from 100, the remainder multiplied first by 0.00287, then by 20, and divided by 3, will give the percentage by weight of the alkaloids of Belladonna Root present in the Plaster

The process of assay adopted by the *U.S.P.* is essentially as follows.—A weighed quantity of 10 grammes of the Plaster in strips is macerated with a mixture of 50 c c of Chloroform and 3 c c of Ammonia Water, the mixture being stirred until the Plaster is completely removed from the cloth, when the Chloroform is transferred to another vessel, the cloth being washed with a mixture of 25 c c of Chloroform and 1 c c of Ammonia Water, if necessary with a further quantity of 25 c c of Chloroform, the chloroformic solutions in each case being removed and added to the main quantity The cloth is dried at a low temperature, cooled and weighed, its weight being deducted from the original weight of Plaster A measured quantity of Alcohol (94.9 p c) equivalent to four fifths of its volume is added to the mixed chloroformic solution, the liquid is gently stirred and allowed to rest until the rubber separates The supernatant liquid is transferred to a separator and agitated for two minutes with 20 c c of a solution prepared by diluting 40 c c of Normal Volumetric Sulphuric Acid Solution with 60 c c of Water After separation, the chloroformic liquid is drawn off and again shaken for two minutes with 10 c c of an acid solution of the same strength, the separated acid liquid mixed with the main quantity, the treatment with this acid solution continued until the shakings fail to give a reaction with Mayer's reagent (Mercuric Potassium Iodide Test Solution) Sufficient Ammonia Water is added to the mixed aqueous acid liquids to ensure an alkaline reaction, and the liberated alkaloids are removed by shaking with Chloroform, using first 25 c c, then 15 c c, and finally 10 c c The chloroformic solutions are separated, mixed, and the Chloroform evaporated off on a water-bath A slight excess of Deci-normal Volumetric Sulphuric Acid Solution is added to the alkaloidal residue, the actual amount added being carefully noted, 10 drops of Chloroform are added, the liquids rotated, and the Chloroform evaporated by means of a water-bath The excess of acid is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, using Cochineal Test-solution as an indicator of neutrality The number of c c of Fiftieth normal Volumetric Potassium Hydroxide Solution used divided by 5, the quotient subtracted from the number of c c of Deci-normal Volumetric Sulphuric Acid Solution first used, divided by the number of grammes of Plaster separated from the cloth, multiplied first by 0.0287 and then by 100, gives the percentage of mydriatic alkaloids present in the Plaster. The *U.S.P.* requires

that it should contain not less than 0.38 p.c. nor more than 0.42 p.c. by weight of such alkaloids

EXTRACTUM BELLADONNÆ ALCOHOLICUM ALCOHOLIC
EXTRACT OF BELLADONNA

Prepared from the Liquid Extract of Belladonna, and readjusted by means of Milk Sugar, so as to contain 1 p.c. of alkaloids

Dose — $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Foreign Pharmacopœias compared under Extractum Vinde

EXTRACTUM BELLADONNÆ LIQUIDUM LIQUID EXTRACT
OF BELLADONNA

A fluid prepared from Belladonna Root in No. 20 powder, with a mixture of 7 volumes of Alcohol (90 p.c.) and 1 volume of Distilled Water, by repercolation, and standardised to contain 0.75 p.c. of alkaloids = $\frac{3}{4}$ grain in 110 minims

A standardised Liquid Extract of Belladonna is official in the *B.P.*, the *U.S.P.* Fluid Extract is prepared with the official standard Belladonna Root in No. 60 powder and contains 0.1 p.c. w/v of mydriatic alkaloids, the *B.P.* from a root which is not standardised, in No. 20 powder. The *P.G.* does not include a fluid extract

Commercial samples of this liquid extract vary enormously in colour and consistence, consequently all preparations made from the liquid extract will also have a tendency to vary in colour, and may necessitate explanations to physicians and patients

Tests.—Liquid Extract of Belladonna has a sp. gr. of from 0.896 to 1.022, and may contain from 6.2 to 26.2 p.c. w/v of total solids and about 70 p.c. w/v of Absolute Alcohol. The *B.P.* process of assay consists in extracting the alkaloids from 10 c.c. of the liquid extract diluted with 5 times its volume of Water, by adding a decided excess of Ammonia Solution, and shaking out with three successive quantities each of 10 c.c. of Chloroform. The chloroformic layers are separated in each instance and the alkaloids are removed from the mixed chloroformic solutions by shaking with a mixture of 5 c.c. of Diluted Sulphuric Acid mixed with twice its volume of warm Water, the operation being repeated to ensure their complete extraction from the Chloroform. The mixed acid liquids are shaken with a small quantity (3 c.c.) of Chloroform, the latter separated and rejected, an excess of Ammonia Solution is added, and the liberated alkaloids are shaken out with two successive quantities each of 10 c.c. of Chloroform. The mixed chloroformic solutions, after being shaken with a little Water (5 c.c.) made faintly alkaline by the addition of one drop of Ammonia Solution, are separated and evaporated, the residue being subsequently weighed after drying at a temperature under 100° C. (212° F.). The actual amount of alkaloid present is then determined volumetrically by dissolving this residue in 10 c.c. of Deci-normal Volumetric Hydrochloric Acid Solution, the excess of acid being determined by titration with Centi-normal Volumetric Sodium Hydroxide Solution, Tincture of Cochineal being used as an indicator of neutrality. 10 c.c. of Deci-normal Volumetric

Hydrochloric Acid Solution being equivalent to 100 c.c. of Centi-normal Volumetric Hydrochloric Acid Solution, if the number of c.c. of Centi-normal Volumetric Sodium Hydroxide Solution required to restore neutrality be deducted from 100 and the product multiplied first by 0.00287 and then by 10, the result will be the weight in grammes of alkaloids present in 100 c.c. of the Fluid Extract. The *B.P.* requires 0.75 p.c. w/v of alkaloids. This process has been subjected to prolonged and severe criticism, and numerous processes have been suggested to fill its place. The obstinate emulsions formed in the very first stages of the process are the worst features. Of the many modifications suggested for the removal of the defects attending the *B.P.* method that of Bird's is probably the most generally used. It is rapid in execution, is free from emulsification, it extracts practically the whole of the alkaloids in a pure condition from any sample of fluid extract, and the results are about from 5 to 6 p.c. higher than by the *B.P.* process. A measured quantity of 10 c.c. is mixed with 2 c.c. strong Ammonia Solution and shaken vigorously with 16 c.c. of a mixture of 3 volumes of Amylic Alcohol, 1 volume of Chloroform and 4 volumes of Ether. The ethereal layer is separated, washed in a second separator with 4 c.c. of Water, added in two portions. The washings are returned to the first separator and the mixed fluids again shaken with 8 c.c. of the Amylic-chloroform ether mixture. The ethereal layer is transferred to the second separator and again washed with 1 c.c. of Water, separated and returned to the first separator. It is extracted a third and a fourth time with 7 c.c. of the above-mentioned mixture and washed in the second separator with 1 c.c. of Water. The mixed ethereal liquids are now extracted four times, first with a mixture of 4 c.c. of Normal Volumetric Sulphuric Acid Solution and 6 c.c. of Water, and then three times in succession with 3 c.c. of Water. Sufficient Ammonia Solution is added to the mixed acid liquids to render them distinctly alkaline in reaction, and they are shaken out four times with Chloroform, using first 10 c.c. and then three successive quantities of 5 c.c. The mixed chloroformic solutions are evaporated on a water-bath at a temperature below 100° C (212° F) to a constant weight, and titrated in the usual manner. Bird has pointed out (*P.J.* [4] 8, 432) that Methyl Orange Solution gives good results as an indicator of neutrality for Belladonna alkaloids. He states that the difference between the results by weight and titration should not exceed 3 or 4 p.c.

The method adopted by the *U.S.P.* is essentially as follows — A measured quantity of 10 c.c. of the fluid extract is diluted with an equal volume of Water and shaken with 20 c.c. of Chloroform and 2 c.c. of Ammonia Solution. The chloroformic layer is removed to a second separator, the treatment being continued with two portions each of 10 c.c. of Chloroform. The mixed chloroformic solutions are now shaken for one minute with 8 c.c. of Normal Volumetric Sulphuric Acid Solution and 20 c.c. of Water. The chloroformic layer is removed and rejected, the aqueous acid liquid is filtered into a clean separator, the first separator and filter being washed with 10 c.c. of Water and

the washings added to the main quantity. After the addition of 4 c c of Armonia solution the latter is well shaken with 20 c c of Chloroform, the chloroformic layer is separated and the extraction repeated with two separate portions of 10 c c of Chloroform. The mixed chloroformic solutions are evaporated on a water-bath and the residue heated till perfectly dry. The residue is then dissolved in 5 c c of Tenth-normal Volumetric Sulphuric Acid Solution, and the excess of acid titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, 5 drops of Cochineal or of Iodeosin Test-solution being employed as an indicator of neutrality. If the number of c.c. of Fiftieth-normal Volumetric Potassium Hydroxide Solution required to neutralise the excess of acid be divided by 5, the quotient subtracted from 5, and then multiplied by 0.0287 and by 10, the product will be the weight in grams of mydriatic alkaloids contained in 100 c c of the Fluid Extract. The opening stages of the *USP* process are very similar to those of the *BP*, and it has been remarked (*P.J.* '07, 1, 393) that it is rather surprising that, after the somewhat harsh criticism to which the process of the *BP* has been subjected, the revisers of the *USP* should adopt a similar one, but on the whole comparatively little difficulty was experienced with the *USP* process, and subsequent experience has shown that if carried out to the letter sharp separations will be obtained. It must not, however, be too hastily assumed that the process is faultless and it is finally concluded that Bud's modification of the *BP* process is more suited to the requirements of pharmacists.

In the author's laboratory the *USP* process has given satisfactory results and is certainly considered preferable to the *BP* process. No emulsification occurred in the earlier stages of the process, the extracted alkaloid was of good appearance and free from dark colour. A note specifically instructed to do so in the *USP* monograph, and these were in nearly all cases weighed when constant, and the difference between the gravimetric and the volumetric estimations never amounted to more than from 3 to 4 p c.

LINIMENTUM BELLADONNÆ. LINIMENT OF BELLADONNA.

Liquid Extract of Belladonna, 10, Camphor, 1, Distilled Water, 2, Alcohol (90 p c), *q.s.* to yield 20

Prescribing Notes—*Prescribed with equal parts of Soap Liniment or Compound Camphor Liniment. Does not mix readily with fixed Oils. When an oily liniment is required is better to order the Chloroform of Belladonna mixed with Olive or Almond Oil.*

Foreign Pharmacopœias—Official in U.S., Camphor 1, Fluid Extract of Belladonna to make 20, Mex. (Acete de Belladonna), Dried Leaves 1, Sesame Oil 10. Not in the others.

SUPPOSITORIA BELLADONNÆ. BELLADONNA SUPPOSITORIES

Made with alcoholic Extract of Belladonna and Oil of Theobroma. Each Suppository contains $1\frac{1}{2}$ grain of Extract, = about $\frac{1}{2}$ grain of alkaloid.

TINCTURA BELLADONNÆ. TINCTURE OF BELLADONNA

1 of Liquid Extract of Belladonna diluted with Alcohol (60 p c) to yield 15

Dose—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Span, Swiss and U S, 1 in 10, Ital, Mex and Port, 1 in 5, Russ, 1 in 12 Port, also 1 in 1, from leaves. All by weight except U S. Not in the others.

The Brussels Conference (1906) adopts a strength of 10 p c, and prepares it by percolation with Alcohol (70 p c). The Belg, Fr and Swiss Ph adopt this standard.

Tests—Tincture of Belladonna, *BP*, is prepared from the standardised Fluid Extract and is officially required to yield not less than 0.048 p c nor more than 0.052 p c w/v of alkaloid. The word 'alkaloid' is presumably intended to read 'alkaloids'. The Fluid Extract from which it is prepared is officially required to yield a fixed percentage of alkaloids. A measured quantity of 100 c c is employed in the process of assay. The Alcohol is removed by evaporating to a low bulk and the resultant product is assayed by the process described under *Extractum Belladonnæ Liquidum*. In calculating the result of the titration the number of c c of Centi-normal Volumetric Sodium Hydroxide Solution used should be deducted from 100 and the product multiplied by 0.00287, which will give the weight in grammes of alkaloids present in 100 c c of the Tincture.

The *USP* evaporates 100 c c of the Tincture until it is reduced to about one-tenth of its volume, adds sufficient Alcohol (94.9 p c) to redissolve any substance which has separated out and employs the process mentioned under *Extractum Belladonnæ Fluidum*. The number of c c of Fiftieth-normal Volumetric Potassium Hydroxide Solution required to restore neutrality divided by 5, subtracted from 5, and multiplied by 0.0287, gives the weight in grammes of mydriatic alkaloids present in 100 c c of the Tincture, it should be 0.3 p c w/v.

The specific gravity of the Tincture varies according to the specific gravity of the liquid extract with which it is prepared, it is usually about 0.910, it contains about 1 p c w/v of total solids, and about 60 p c w/v of Absolute Alcohol.

UNGVENTUM BELLADONNÆ. BELLADONNA OINTMENT

8 of Liquid Extract of Belladonna evaporated to 1, and mixed with 9 of Benzoated Lard.

Contains 0.6 p c of Alkaloid.

Foreign Pharmacopœias—Official in Belg and U S, Extract 1 in 10, Fr (*Pommade Belladonnée*), Extract 3, Glycerin 2, Benzoated Lard 25, Ital (*Pomata di Belladonna*), Extract 10, Glycerin 5, Benzoated Lard 85, Mex (*Pomada de extracto de Belladonna*), Extract 1, Lard 7½, Port (*Pomada de Belladonna*), aqueous Extract 1, Lard 9, (*Pomada de Belladonna Forte*), Alcoholic Extract 1, Lard 9, Span (*Pomada de Belladonna*), Extract 1½ in 10. Not in the others.

Not Official.

CHLOROFORMUM BELLADONNÆ—Belladonna Root, in powder, 20, percolate with Chloroform *q s* to yield 20.—*Squire's Companion* 1864.

Applied with equal parts of Camphor Liniment or Olive Oil, for painful rheumatism

It is well known that this preparation only extracts about half of the total alkaloids. By mixing the Root (in No 40 powder) with Slaked Lime and powdered Carbonate of Ammonium, four-fifths of the alkaloid will appear in the first 1 in 1 percolate. —Squire's *Chemistry*, 1894

Belladonna Root, 1 No 60 powder, 100, Solution of Ammonia, 25 Absolute Alcohol *q s*, Chloroform *q s*. Moisten the Belladonna root with the solution of Ammonia and set aside for 24 hours. Transfer to a 1-litre percolator with a menstruum consisting of 1 of Absolute Alcohol to 7 of Chloroform until 100 of percolate is obtained —*B P C*

The above formula is the outcome of a series of experiments undertaken by R. Wright, full particulars will be found in *Y B P* '03, 589; *P J* '03, 153; *C D* '03, 246; *Y B P* '07, 367, 371; *P J* '07, 106; *C D* '07, 171

Samples of Chloroform of Belladonna were prepared by the *B P C* process, and by that recommended by the *B P C*. That prepared by the *Com pamon* 1894 process had a specific gravity of 1.476, it contained 3.11 p.c. w/v of total solids, and when assayed according to an adaptation of the *U S P* process for the assay of Belladonna Root yielded gravimetrically 0.25 p.c. w/v of alkaloids and volumetrically 0.236 p.c. w/v of alkaloids, calculated as Atropine. When assayed by the process recommended by Farr and Wright it yielded gravimetrically 0.24 p.c. w/v of alkaloids and volumetrically 0.235 p.c. w/v of alkaloids, calculated as Atropine. This latter process gave no trouble in its manipulation, and emulsions were produced which could be separated even on standing, and clear separations had to be produced mechanically. The product of the *U S P* process of assay worked admirably, it yielded the alkaloids practically free from colour and in a pure condition, no trouble was experienced through emulsification. The *B P C* product had a specific gravity of 1.419, it contained 3.79 p.c. w/v of total solids, and showed when assayed by a method founded on a modification of the *U S P* process for the assay of Belladonna Root, gravimetrically 0.38 p.c. w/v of alkaloids, volumetrically 0.365 p.c. w/v of alkaloids, calculated as Atropine. By the process recommended by Farr and Wright it indicated gravimetrically 0.22 p.c. w/v of alkaloids, and volumetrically 0.209 p.c. w/v of alkaloids, calculated as Atropine.

COLLODIUM BELLADONNÆ *Syn.* ENIPLASTRUM BELLADONNÆ FLUIDUM —Liquor Extract of Belladonna, 50, Canada Turpentine, 1 Castor Oil, 2 and Ether, 40. Mix and digest for 12 hours, filter and add Camphor, 15, Pyroxilin 25, and Ether (sp. gr. 0.720), *q s* to make 100 —*B P C*

This is a modification of the formula given in *B P C* *Formulary* 1901

GLYCERINUM BELLADONNÆ —Green Extract of Belladonna, 8, boiling Distilled Water 1, Glycerin, to 16 —*B P C* *Formulary* 1901

This has been incorporated in the *B P C*

This is practically the same as the *U S P* formula, it also appears in other Hospital Pharmacopœias, with varying quantities.

Official in Part, 1 Extract in 10

LINIMENTUM BELLADONNÆ COMP —Liniment of Belladonna, 7, Chloroform of Belladonna, 1, mix. For application to the loins in lumbago it should be sprinkled on impermeable piline (not *gummi piline*), and firmly pressed with the hands on the part for five minutes to ensure perfect contact, it should then be kept on for at least 16 or 12 hours.

Peter Squire, who suffered much from lumbago, found this more effectual and much more convenient than Belladonna Plasters.

LINIMENTUM BELLADONNÆ CUM CHLOROFORMO —Chloroform, 12 50, Liniment of Belladonna, *q s* to make 100 —*B P C*

ETHEREAL TINCTURE OF BELLADONNA (*Squibb's*) —Substituted by the Ether for Rectified Spirit in the Liniment of *B P* '55 —*J.* '90, 11 67

Not Official BENZIN

PETROLEUM BENZIN PETROLLEUM ETHER

A purified distillate from American Petroleum. It is a transparent, colourless, highly inflammable liquid, possessing a characteristic odour.

It should be preserved in well-stoppered bottles and in a cool atmosphere. It forms a highly explosive mixture with air.

Solubility—Insoluble in Water, about 1 in 6 of Alcohol (90 p c), readily soluble in Ether, Chloroform, fixed and volatile Oils.

Medicinal Properties—Used in seborrhoea, in acne, and generally for the purpose of dissolving off grease from the skin. It is highly inflammable, and must not be used near a fire or naked flame.

Foreign Pharmacopœias—Official in Ger, Jap, Russ, Swiss and U S. U S has also Benzinum Purificatum.

Tests—The distinguishing tests for Benzin are its peculiar odour, the specific gravity which should be 0.670 to 0.675, the boiling point which should be from 45° to 60° C (113° to 140° F), and that it does not produce the odour of Nitro benzene when treated with ten times its volume of a mixture of 1 volume of Sulphuric Acid and 1 volume of Nitric Acid.

Petroleum Ether sp gr 0.716, generally known under the name of **Petrol**, is used extensively for the internal combustion engines of motor vehicles.

BENZINUM PURIFICATUM—Potassium Permanganate, 1, Sodium Hydroxide, 0.2, Sulphuric Acid, 6, Petroleum Benzin, 100, Water *q s*. Add the Acid to 55 of Distilled Water and when cold pour into a bottle having the capacity of about 200. Add 0.8 of Potassium Permanganate, agitate till dissolved, then add the Petroleum Benzin in four portions, shaking the liquid after each addition, allow the liquids to remain in contact for 24 hours, shaking the bottle at frequent intervals, then decant the Petroleum Benzin, and having dissolved 0.2 gramme of Potassium Permanganate in 24 of Water, in which the Sodium Hydroxide has previously been dissolved, mix and agitate frequently, then decant, repeat the washing with Water, and again decant the Petroleum Benzin.—U S P.

This has been incorporated in the *B P C*.

BENZONINUM.

BENZONIN

FR, BENJOIN, GER, BENZONHARZ, ITAL, BENZOINO, SPAN, BENJUI

A balsamic Resin, obtained from *Styrax Benzoin*, Dry, and probably from other species of *Styrax*, both Siam and Sumatra Benzoin are specifically mentioned in *B P*, but the latter very seldom complies with the official characters.

Siam Benzoin consists of about 38.0 p c of Benzoic Acid, 56 to 57 p c of the Benzoic Ester of Siareosinotannol, and about 5 p c of the Benzoic Ester of Benzoresinol, a small amount of Vanillin, and an oily neutral liquid consisting of the Benzoic Ester of Cinnamyl or Benzyl Alcohol. Sumatra Benzoin consists chiefly of the Cinnamic Ester of Benzoresinotannol, some Cinnamic Ester of Benzoresinol, Styracin, Cinnamic Acid Ester of Phenylpropyl Alcohol, a little Vanillin, free Benzoic and Cinnamic Acids, and traces of Benzaldehyde and Benzol.

Solubility.—The tears are as a rule wholly soluble 1 in 5 of Alcohol (90 p c), 1 in 1 of Ether, and in Solution of Potassium

Hydriodic. The mass contains impurities, which are left after treating it with Alcohol. The Solution in Alcohol or Ether is acid.

B.P. requires Benzoin to be almost entirely soluble in Alcohol (90 p.c.), but Sumatra Benzoin is rarely so.

Medicinal Properties.—Expectorant, styptic, antiseptic, used in making aromatic fumigating pastilles. The **compound tincture** is given internally for chronic bronchitis, the **vapour** or **spray** is used in chronic laryngeal and bronchial catarrh to check abundant secretion and cough, lint soaked in the compound tincture forms a styptic and antiseptic dressing for wounds.

Prescribing Notes.—*If given in the form of mixture the Tincture should be emulsified with Mucilage of Gum Acacia, or yolk of Egg. A nice lotion to protect the face from the heat of the sun is made with Tincture of Benzoin 1, Rose Water 40.*

Official Preparation.—*Tinctura Benzoini Composita.* Used in the preparation of Acidum Benzoicum, Adeps Benzoatus, and Unguentum Cetacei.

Not Official.—*Tinctura Benzoini, Insufflatio Benzoini, Lint Virginal, Lotic Benzoini, Sevum Benzoatum, Sobum Benzoinatum, Unguentum Benzoini, Vapor Benzoini.*

Foreign Pharmacopœias.—Official in Austri, Belg, Dan, Dutch, Fr, Ger (Benzoe), Hung, Ital, Jap, Norw, Port, Russ, Mex, and Span., Swed, Swiss and U.S.

Descriptive Notes.—Several varieties of Benzoin are met with in commerce known respectively as Siam, Saigon, Sumatra, Penang and Palembang. The last three kinds are produced in different districts of Sumatra, probably by different trees. Of each kind several grades occur, varying in freedom from foreign matter and in their appearance. Siam Benzoin almost always occurs in more or less distinct tears, or when broken into masses they leave interstices and the masses have a translucent or varnished surface. The odour of Siam Benzoin recalls that of Vanilla. The tears may vary in size from that of small shot to an inch or two (25 to 50 mm) in length and breadth and $\frac{1}{2}$ inch (12.5 mm) in thickness, the tears being usually flattened and of a pale brown externally but milky-white internally. It is remarkably free from impurity and rich in Benzoic Acid, and contains Vanillin. Sumatra Benzoin occurs in solid masses, presenting irregular white tears immersed in a dull greyish-brown resin, and does not exhibit the translucent varnished appearance of the Siam lump Benzoin. It has a characteristic odour resembling Storax rather than Vanilla. It contains Cinnamic as well as Benzoic Acid. It is chiefly produced in Sumatra. Penang Benzoin resembles that of Sumatra, but has a much more pronounced Storax or Hyacinth odour. It is probably produced in W. Sumatra from *Styrax subdenticulatum*, Miq., and comes *via* Penang. It is rarely met with in the form of tears. Saigon Benzoin, imported from Cochinchina, resembles that of Sumatra in appearance and odour. Palembang Benzoin is distinguished by almost entire absence of white tears and a varnished, not dull appearance as in Sumatra Benzoin, it has but little odour. It is produced in the East of Sumatra.

Unfortunately the commercial Benzoin which has a Storax odour is sometimes sold under the name of Penang, and sometimes sold under that of Sumatra Benzoin, so that one is often mistaken for the other. Cinnamic Acid is present in Sumatra Benzoin and the Storax smelling Benzoin, but absent in Siam, Saigon, and Palembang Benzoin, according to E. Wightman Bell, who states that Siam is the richest in Benzoic Acid, yielding 30 to 37 p c.

Tests—The distinguishing tests for Benzoin are its physical appearance and agreeable aromatic odour, that it readily softens when warmed, and on being subjected to a still higher degree of heat yields vapours of Benzoic Acid, it should be almost entirely soluble in Alcohol (90 p c) and in Potassium Hydroxide Solution. The Acid and Saponification values afford useful data for judging the quality of the Gum, but no mention of these values is made in either the *BP*, *USP*, or *PG*. The Acid value of a good specimen of Siam Benzoin should be between 140–170, and the Saponification value 220–240. Three good commercial samples of Siam Benzoin examined in the author's laboratory, leaving only about 1 p c insoluble in Alcohol (90 p c), gave Acid values ranging from 154.6 to 184.8, and Saponification values ranging from 197 to 218. A sample of Siam Benzoin which left 14.7 p c insoluble in Alcohol (90 p c) yielded an Acid value of 123.2, and a Saponification value of 184.8. A fine commercial sample of Sumatra Benzoin, leaving 1.4 p c insoluble in Alcohol (90 p c), had an Acid value of 132.2, and a Saponification value of 181.5. The five samples of Siam Benzoin left ashes ranging from 0.01 p c to 0.88 p c, the Sumatra Benzoin left 0.68 p c of ash. The solubility in Alcohol (90 p c) varies with the origin of the Gum. The best Siam Benzoin is as a rule wholly soluble 1 in 5. Sumatra Benzoin, although distinctly specified, would appear from the description not to be intended for use, since it is almost impossible to obtain it in commerce with less than 7 to 10 p c of residue, which is not presumably covered by the words 'almost entirely soluble in Alcohol (90 p c)'. In view of the use of Benzoin in the preparation of the Compound Tincture, the question of the solubility of the sample assumes some importance.

Attfield has pointed out in his *Digest of Researches and Criticisms* (Report for 1898) that Benzoin containing the usual varying proportions of bark (1 to 30 p c) may be employed in preparing the Compound Tincture, due allowance being made for the insoluble matter, until it can be shown that the attached bark has parted with harmful soluble matter to the Alcohol (90 p c). The *USP* requires Benzoin to be almost wholly soluble 1 in 5 parts of warm Alcohol (94.9 p c), the *PG*, which only recognises the Siam variety, requires that it shall leave not more than 5 p c by weight of insoluble residue when exhausted with boiling Alcohol (90 p c).

The more generally occurring adulterants are inferior varieties of Gum, Colophony, Storax, Turpentine, and mineral matter. The *BP* does not include tests for any of these substances. The *USP* and *PG* require that it should not, on incineration, leave more than 2 p c by weight of Ash. The *PG* also requires that crystals of Benzoic

Acid shall separate from the colourless fluid obtained on warming 1 part of the Gum with 10 parts of Carbon Bisulphide Styra^x lowers the Acid value, and Turpentine the Ester and Saponification value

Preparation

TINCTURA BENZOINI COMPOSITA. COMPOUND TINCTURE OF BENZOIN.

B P Syn —FRIAR'S BALSAM *N O Syn* —TRAUMATIC BALSAM

Benzoin, 8, prepared Storax, 6, Balsam of Tolu, 2, Socotrine Aloes, $1\frac{1}{2}$ (less $\frac{1}{4}$)^{*}, macerated with Alcohol (90 p c) to yield 80. (1 in 10)

Although Sumatra Benzoin is permitted by the Pharmacopœia only Siam should be used, on account of its superior solubility

Dose.— $\frac{1}{2}$ to 1 fl drim = 18 to 36 cc

Foreign Pharmacopœias.—Official in Mex (Tinctura de benjuí compuesta), Port, Swed, and U.S., Fr (Teinture Balsamique), the tinctures vary considerably in composition and strength Not in the others

Benzoin, in No 40 powder, 10, Purified Aloes, in No 40 powder, 2, Storax, 8, Balsam of Tolu, 4, Alcohol (95 p c), *qs* to make 100 —*U.S.P.*

Tests.—It possesses a specific gravity of about 0.900, and contains from 17 to 18 p c of total solids and about 75 p c w/v of Absolute Alcohol It has been suggested (C.D. '02, 1432) that independently of the determination of extractive matter, a determination of free and combined Cinnamic Acids should be made, and that a good Tincture should yield not less than about 5 p c of balsamic acids calculated as Benzoic Acid, of which neither more nor less than two-fifths should be present in an uncombined condition

Not Official

LOTIO BENZOINI—A nice lotion to protect the face from the sun is made with Tincture of Benzoin, 1, Rose Water, 40 —*Squire*

This has been incorporated in the *B.P.C.*

LAIT VIRGINAL—Tincture of Benzoin, 2 fl drim, Rose Water, to 8 fl oz

A proof-spirit tincture gives the best result, but the Milk is greatly improved by the addition of 3 fl drim of Glycerin to the Water Orange-flower Water or other aromatic Water may also be used —*Pharm. Form.*

TINCTURA BENZOINI.—1 of Benzoin in powder, macerated with Alcohol (90 p c) *qs* to yield 10

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr., Ger., Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U.S., 1 in 5, all by weight, except U.S. Swiss includes also Tinct Benzoës Æthereæ 1 in 5 The Austr and Belg Ph require the tincture to yield at least 18 p.c. of dry residue, the Dutch not less than 15 p c

INSUFFLATIO BENZOINI (*Vigier*)—Tincture of Benzoin, 1, Boic Acid, 1, Starch Powder, 1 Mix, and let the Alcohol evaporate. Used as a snuff in coryza —*T.G.* '88, 141

SEVUM BENZOATUM.—Benzoated Suet is prepared in the same manner as Benzoated Lard, Prepared Suet being used in place of Lard —*Ind* and *Col. Add*

^{*} To be exact, 18 grains are to be taken from every $1\frac{1}{4}$ oz of Aloes

Sebum Benzoinatum —Suet, 100, Benzoin, 4, Dried Sulphate of Soda, 6
Swiss

UNGUENTUM BENZOINI —Benzoin, in fine powder, 1, Adeps, 4

VAPOR BENZOINI —Compound Tincture of Benzoin, 60 minims in a pint of Water at 140° for each inhalation

For bronchitis and laryngitis

This has been incorporated in the *B P C* as follows —

Compound Tincture of Benzoin, 0 50, Water at 60 °C, 100

BENZOL.

A colourless, inflammable, volatile liquid, containing about 70 p c of Benzene and 20 to 30 p c of Toluene. It is obtained from Coal-tar Naphtha and must not be confused with Benzoin from Petroleum. See p 237

Introduced into *B P* as a solvent for india rubber

Solubility.—Insoluble in Water. Soluble in all proportions of Absolute Alcohol, Chloroform, and Ether

Medicinal Properties —Stated to be useful in influenza

Only the purest crystallisable Benzol should be used for internal administration

Dose —For children 3 minims, adults 5 minims, prescribed in capsules every two or three hours, or in mixture

Benzol 80 minims, Alcohol (90 p c) $\frac{1}{2}$ oz, Sp Chloroform 3 fl drms, Mucilag Trag to 8 oz, dose, $\frac{1}{2}$ oz every three hours, in lemonade —*B M J* '92, i 171, '93, ii 1425, *L* '92, i 234

Foreign Pharmacopœias —Official in Dutch, Fr (Benzine), Mex (Bencina), Port (Benzina), and Swed

Tests —The distinguishing tests for Benzol are its peculiar somewhat aromatic odour, its specific gravity, about 0 885. It should commence to boil at 80° C (176° F), nine-tenths should distil below 100° C (212° F), and the remainder below 120° C (248° F)

The *U S P* gives the specific gravity as 0 871 at 25° C (77° F), and the boiling point as 80 4° C (176 7° F)

The more generally occurring impurities are readily oxidisable organic compounds or Thiophene. The *B P* does not include tests for these compounds. The *U S P* requires that Sulphuric Acid mixed with an equal volume of Benzol shall not become coloured, indicating the absence of readily charred organic impurities, and that no green or blue tint shall be developed on shaking it with quarter its volume of Sulphuric Acid and one drop of Fuming Nitric Acid, indicating the absence of Thiophene

Not Official

BERBERIS

The Bark of the root of *Berberis vulgaris*, L

It contains the alkaloids, Berberine $C_{22}H_{17}NO_4$, and Oxyacanthine $C_{11}H_9NO_2$

The dried stem of *Berberis Aristata* is official in the *Ind* and *Col Add* for India and the Eastern Colonies, also **Tinctura Berberidis**, 1 in 10 of Alcohol (60 p c), dose, 30 to 60 minims = 1.8 to 3.6 c c

Medicinal Properties—A bitter tonic. Has been used with success in intermittent fevers.

It has also been used in India as a local application in affections of the eye.

EXTRACTUM BERBERIDIS FLUIDUM.—Made with Alcohol (60 p c) One fl. oz. of Extract is equal to 1 oz. of Bark.

Dose—20 to 60 minims = 1.2 to 3.6 c c

Liquor Berberidis Concentratus (1 in 2), dose 30 to 60 minims, is official in the *Ind* and *Col Add* for India and the Eastern Colonies.

BERBERINÆ PHOSPHAS—This is the most soluble salt of Berberin. Soluble 1 in 15 of Water, 1 in 9 of hot Water, but part separates out on standing, it is also thrown down as a yellow precipitate by excess of Alcohol.

Dose.—1 to 5 grains = 0.06 to 0.32 gramme

Not Official

BETEL.

The Leaves of *Piper Betle*, L., are official in the *Ind* and *Col Add* for India and the Eastern Colonies. It is largely employed in India as a masticatory in conjunction with Lime and the nut of *Azeca Catechu*, L.

Not Official.

BETULÆ ALBÆ OLEUM.

BIRCH TAR OIL.

Syn—**OLFUM RUSCI**

A bituminous liquid obtained by destructive distillation of the Wood of *Betula alba*, L., produced in Russia.

Principally employed as an application in skin diseases, also in rheumatism and gout. Was at one time given also internally, in doses of 3 to 8 grains = 0.2 to 0.52 gramme, in pill.

Official in Austr., Jap. and Swiss.

Oleum Betulinum Rectificatum—A light-brown Oil, obtained by the steam distillation of Birch Tar Oil.

The active constituents of the Rectified Oil are probably Guaiacol and Cresol—*P J* (3) 661.

Solubility—Almost insoluble in Water, soluble in all proportions of Alcohol, Chloroform or Ether.

Official in Dutch.

Tests—The specific gravity should be from 0.900 to 0.920. A saturated aqueous solution of the Oil gives a green coloration on the addition of a few drops of a 1 in 1000 aqueous Ferric Chloride Solution.

Not Official

TINCTURA RUSCI (*Hebra*)—Birch Tar Oil, 25; Oils of Lavender, Rue and Rosemary, of each 1, Ether, 86, Alcohol (50 p c) 36.

Official in Austr., Lavender Oil, 1, Rosemary Oil, 1, Birch Tar Oil, 26, Ether, 86, Alcohol, 86.

UNGUENTUM OLEI BETULÆ.—Birch Tar Oil, 5 fl. drms., Yellow Beeswax, 120 grains, melt the Beeswax, add the Oil, and stir till cold.

* Used in psoriasis and dry eczema.

Caution.—The use of this Ointment in eczema demands care.

UNGUENTUM BETULÆ COMPOSITUM — Oil of Cade, 10, Resorcin, 1, Ichthyol, 1, Birch Tar Oil, 1, Lard, 30 — *St George's*

Not Official

BISMUTHUM

B1, eq 207 30

FR, BISMUTH PURIFIC, GER, WISMUT, ITAL, BISMUTO, SIAM, BISMUTO

Bismuth in its crude state is generally impure the official salts, however, are required to give no reaction with a special test for Selenium and Tellurium

Official in Mex (Bismuto), Port, Span and Swiss

Official Bismuth Salts — Bismuthi Carbonas, Bismuthi Oxidum, Bismuthi Salicylas, and Bismuthi Subnitras

Not Official — Bismone (Colloidal Bismuth Oxide), Bismuthum Purificatum, Bismuthi Benzoas, Bismuthi Betanaphtholas (Oiphol), Bismuthi et Cerii Salicylas, Bismuthi Citras, Bismuthi et Ammonii Citras, Bismuthi et Cinchonidine Iodidum (Erythrol), Bismuthi Di-thio Salicylas (Thioform), Bismuthi Iodoresorcin Sulphonas (Anusol), Bismuthi Methylenedigallas (Bismal), Bismuthi Nitras, Bismuthi Oleas, Bismuthi Oxichloridum (Pearl white), Bismuthi Oxydodogallas (Ainol), Bismuthi Phenolas, Bismuthi Phosphas, Bismuthi Quinolini Sulphocyanidum (Crurin), Bismuthi Sulphis, Bismuthi Subgallas (Dermatol), Bismuthi Subiodidum, and Bismuthi Tribromophenolas (Xeroform)

BISMUTHI CARBONAS.

BISMUTH OXYCARBONATE

($\text{Bi}_2\text{O}_3\text{CO}_3$), H_2O , eq 1029 70

FR, SOUSCARBONATE DE BISMUTH, GER, WISMUTSUBCARBONAT, ITAL., BISMUTO CARBONATO, SPAN, CARBONATO DE BISMUTO

A white or almost white, odourless and tasteless amorphous powder, which varies much in density, the lighter variety is most suited for dispensing, being more easily suspended

It may be prepared from the Subnitrate by precipitation with Ammonium Carbonate

Solubility — Soluble with effervescence in Nitric Acid, insoluble in Water

Medicinal Properties — Similar to the Subnitrate, and often preferred to it

The Carbonate is most generally useful as a gastric sedative, the Subnitrate is the most effective as an intestinal antiseptic, the Salicylate being weaker and the Carbonate inert for this purpose — *L* '05, 1 432

Dose — 5 to 20 grains = 0.32 to 1.3 gramme

Prescribing Notes — *Suspended in mixture by Compound Tragacanth Powder*

Mucilage of Gum Acacia is not a good vehicle for Bismuth salts. On standing, a compact mass forms at the bottom of the bottle, which is difficult to diffuse

When Sodium Bicarbonate is to be given with a Bismuth salt, the Carbonate should be selected

a good one for pyrosis *Bismuthi Carbonatis*, 2
 drms, *Pulv. Tragac.* Comp 1 drms, *Aq. Flor.*
Aurant. (glycerine of each 2 fl drms, *Aquæ Chloroformi*, 1½ fl oz, *Aquam ad* 6
 fl oz 3 to 4 teaspoonfuls three times a day after meals

Official Preparation—*Trochiscus Bismuthi Compositus*

Not Official—*Glycerinum Bismuthi Carbonatis*, *Mistura Bismuthi*,
Mistura Bismuthi cum Soda, *Pastillus Bismuthi*, *Pastillus Bismuthi et Mor-*
phinæ.

Foreign Pharmacopœias—Official in Dutch, Jap., Mex. (*Carbonato*
de Bismuto), Port., Span. and U.S. Not in the others.

Tests.—The distinguishing tests for Bismuth Carbonate are that it dissolves with effervescence in Hydrochloric Acid yielding a solution from which (1) Hydrogen Sulphide throws down a brownish-black precipitate insoluble in Ammonium Hydrosulphide Solution, in Potassium or Sodium Hydroxide Solution, but soluble in hot Nitric Acid, (2) except in the presence of Citric Acid or Citrate, Ammonium, Potassium or Sodium Hydroxide Solution throws down a white precipitate insoluble in excess, (3) the copious dilution with Water of a strong solution of the salt in a sufficiency of mineral acid results in the formation of a white precipitate, in the case of the solution in Nitric Acid if no precipitation results on dilution, the addition of Ammonium or Sodium Chloride Solution brings about immediate precipitation, the presence of Tartaric Acid not affecting the precipitation, (4) the addition of Potassium Chromate Solution causes a yellow precipitate insoluble in Potassium or Sodium Hydroxide Solution, soluble in dilute Nitric Acid. When treated with Hydrochloric Acid it effervesces briskly evolving a gas, which, passed through Calcium Hydroxide Solution, affords a white precipitate. The salt is officially required to indicate 99.93 p.c. of Bismuth Carbonate, as ascertained from the weight of Bismuth Sulphide (99.0 p.c.) resulting from precipitation with Hydrogen Sulphide, the *B.P.* employing the latter reagent for the determination of the Bismuth. A weighed quantity of 1 gramme of the Carbonate is dissolved in a little Hydrochloric Acid, the solution diluted with Water acidified with Hydrochloric Acid, and Hydrogen Sulphide is passed through the solution until the Bismuth is completely precipitated. The precipitate is filtered off, rapidly washed with Water and dried at 100° C. (212° F.) till constant in weight, and when cool, weighed. The weight of Bismuth Sulphide should amount to 0.99 grammic. The *U.S.P.* method of determination is to ignite at a red heat and to weigh the residue of Bismuth Oxide which should amount to not less than 90 p.c., equivalent to 100 p.c. of Bismuth Subcarbonate. The results obtained by the Sulphide method of determination are likely to be much higher than those obtained by the *U.S.P.* method owing to the tendency towards co-precipitation of Sulphur which would not be washed out and which would be weighed as Bismuth Sulphide. It is considered (*C.D.* '98, i. 674, '98, ii. 348) that the differences by the ignition method are much smaller than those occurring in the Sulphide method.

The Carbonate is not official in the *P.U.*

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead, Magnesium, Silver and Zinc, Chlorides, Nitrates

and Sulphates, Selenium and Tellurium. These are characteristically lumped together by the *BP* without any regard to their relative importance, and so long as the salts are 'suitably treated,' the usual tests for these substances are employed. A standard of 2 parts per 1,000,000 is suggested (*CD* '08, i 795) for Arsenic. It should yield no reactions for Arsenic when examined by the Bettendorf's test. When 3 grammes of the salt is dissolved in just sufficient warm Nitric Acid to affect solution, and this solution be then poured into 100 c c of Water, filtered, the filtrate evaporated on a water-bath to 30 c c, and again filtered, portions each of 5 c c of this filtrate should not yield a blue supernatant fluid on the addition of an excess of Ammonia Solution when the precipitate is allowed to settle, indicating the absence of Copper, should not become cloudy when mixed with an equal volume of Diluted Sulphuric Acid, indicating the absence of Lead, should not yield a precipitate on the addition of Hydrochloric Acid, indicating the absence of Silver, nor yield a turbidity on the addition of Barium Chloride Solution, indicating the absence of Sulphates. If 1 gramme of the salt be dissolved in equal parts of Acetic Acid and Water, and the Bismuth be completely removed by Hydrogen Sulphide, the filtrate from the Sulphide precipitate should yield no residue on evaporation, indicating the absence of metals of the alkalis and alkali earths. The *USP* disregards the presence of Iron and Zinc, Selenium and Tellurium, except in so far as the two latter are covered by Bettendorf's Arsenic test. A special test for Ammonium salts with Potassium Hydroxide Solution is given, a limit for Chlorides is adopted and in contradistinction to the *BP* the presence of Nitrates is prohibited, the *BP* allows not more than the slightest reactions. The commercial Carbonate invariably contains more than a trace of Nitrate (*PJ* (3) xvi 936, (3) xviii 721, 780), but it can be obtained in commerce free from Nitrate (*CD* '98, i 837). Nitrates may be detected, if present, by the Ferrous Sulphate and Sulphuric Acid test given under that heading in the small type below, Chlorides by the test given under the heading of Silver Nitrate. When testing for Selenium and Tellurium the *BP* removes the major portion of the Bismuth as an oxy-salt by the addition of Sodium or Ammonium Chloride to the Nitric Acid solution, and adds an excess of Sodium Sulphite to the filtrate, no precipitate or coloration should be given after 12 hours, indicating the absence of Selenium and Tellurium. A delicate test for Tellurium given (*CD* '97, i 631) is to dissolve without heat 10 grains of Bismuth salt in 60 minims of strong Hydrochloric Acid mixed with 60 minims of Water, add 10 grains of Sodium Hypophosphite, an evolution of Nitrous fumes will take place in the case of Subnitrate and of Carbonic Anhydride only if it be Carbonate, but no development of colour or precipitation if the Bismuth salt be pure. If Tellurium be present in very small proportion a black precipitate will form, and if Arsenic be the impurity the precipitate will be brown.

Ferrous Sulphate and Sulphuric Acid.—If a mixture of 0.05 gramme of Bismuth Subcarbonate and 5 c c of equal parts of Water and Ferrous Sulphate be agitated and poured as a layer over 5 c c of Sulphuric Acid (free from

Not a brownish red zone should form at the junction of the liquid, indicating a limit of Subnitrate, *U S P*

Silver Nitrate—If the precipitate (if any) formed by the addition of 0.1 c.c. Tenth-normal Volumetric Silver Nitrate Solution to a solution of 0.3 gramme of Bismuth Subcarbonate in 10 c.c. of Nitric Acid be filtered off the clear filtrate should be unaffected by the further addition of the reagent, indicating a limit of Chlorides, *U S P*

Gravimetric Determination—1 gramme of Bismuth Subcarbonate ignited in a porcelain crucible should yield a residue of not less than 0.9 gramme of Bismuth Oxide, *U S P*

Preparations

TROCHISCUS BISMUTHI COMPOSITUS. COMPOUND BISMUTH LOZENGE

2 grains of Bismuth Oxycarbonate, 2 grains Heavy Magnesium Carbonate, and 4 grains Precipitated Calcium Carbonate in each, with Rose basis

Dose.—1 to 6 lozenges

A modification, known as the **Gastric Antacid Lozenge**, has been recommended by Sir W. Roberts, the Bismuth is omitted and Sodium Chloride added—*B M J '89* ii 874.

Foreign Preparation—Official in Port, $1\frac{1}{2}$ grain of subnitrate in each. Not in

Not Official

GLYCERINUM BISMUTHI CARBONATIS—Bismuth Oxynitrate, 2820 grains, Water, 3 fl oz, Nitric Acid, $1\frac{1}{2}$ fl oz. Dissolve the Bismuth Oxynitrate in the mixture of Water and Nitric Acid and pour into a solution of Ammonium Carbonate $5\frac{1}{2}$ oz in Water 30 fl oz, wash the precipitate by decantation, drain, and mix the residue with Glycerin, *q s* to make 10 fl oz. This preparation contains 1 grain of Bismuth Oxycarbonate in 2 minims.—*St Thomas's*

Bismuth Nitrate, in crystals, 100, Nitric Acid, 15, Ammonium Carbonate, 50, Distilled Water, 360, Glycerin, *q s* to produce 100. The product contains about 50 p.c. of Bismuth Carbonate—*B P C*

The *B P C Supplement* gives the first formula as an alternative method, with the *syn* Bismuth Cream

MISTURA BISMUTHI—Glycerin of Bismuth Carbonate, 30 minims; Water, to 1 fl oz.—*St Thomas's*

This has been incorporated in the *B P C*

MISTURA BISMUTHI CUM SODA—Bismuth Oxycarbonate, 15 grains, Sodium Bicarbonate, 10 grains, Tragacanth, 11 grains, Water, to 1 fl oz. This mixture may be made without Tragacanth but the Bismuth Oxycarbonate subsides more quickly.—*St Thomas's*

Sodium Bicarbonate, 10 grains, Bismuth Mixture, *q s* to produce 1 fl oz.—*B P C*

PASTILLUS BISMUTHI.—Carbonate of Bismuth, 3 grains, 3 minims. Rub together and add the mixture to the melted Tragacanth, 14 grains.—*Throat*.

This has been incorporated in the *B P C*.

PASTILLUS BISMUTHI ET MORPHINÆ—Carbonate of Bismuth, 3 grains, Acetate of Morphine, $\frac{1}{10}$ grain, Glycerin, 3 minims, Glycogelatin, 18 grains.—*Throat*

This has been incorporated in the *B P C*.

BISMUTHI OXIDUM.

BISMUTH OXIDE

 Bi_2O_3 , eq 462 24

A pale yellowish-white, amorphous powder

It may be prepared by the interaction of Bismuth Oxynitrate and Sodium Hydroxide Solution at a boiling heat

Solubility—Insoluble in Water, soluble in Nitric Acid mixed with half its volume of Water**Medicinal Properties**—Similar to the subnitrate**Dose**—5 to 20 grains = 0.32 to 1.3 gramme**Not Official**—Bismuthi Oxidum Hydratum and Cremor Bismuthi**Tests**—Bismuth Oxide should answer the tests distinctive of Bismuth given under the Carbonate. It is officially required to contain 99.68 p.c. of Bismuth Oxide, as gravimetrically determined by conversion into Bismuth Sulphide. 1 gramme of the Oxide should yield 1.1 gramme of the Sulphide. The objections to this method of determination are given under Bismuth Carbonate.

The more generally occurring impurities are such as are also found in the Carbonate and are there discussed. In addition it may contain Bismuth Oxy carbonate or the Oxynitrate, or moisture, in which case there will be an appreciable loss of weight when a weighed quantity is heated to incipient redness. Such diminution in weight is officially prohibited.

Not Official.**BISMUTHI OXIDUM HYDRATUM**—A white amorphous powder, soluble in an excess of Hydrochloric Acid and precipitated again on the addition of Water as Oxychloride. It mixes readily with Water to form a cream.**Official in Fr. and Span.****CREMOR BISMUTHI**—Hydrated Bismuth Oxide, 1, Water, 4. Rub together till smooth.

Under the name 'Intestin' a mixture containing Bismuth Oxide, Benzoic Acid and Naphthalene has been introduced.

BISMUTHI SALICYLAS.

BISMUTH SALICYLATE

 $\text{C}_6\text{H}_4(\text{OH})(\text{COO BiO})$, eq 359 19

A white or nearly white, odourless powder, but also supplied in crystals. It should contain 62 to 64 p.c. of Bismuth Oxide.

It may be prepared by precipitating Bismuth Nitrate with a solution of Sodium Salicylate.

Solubility—Insoluble in Water and Alcohol (90 p.c.)**Medicinal Properties**—An excellent intestinal antiseptic and sedative, has been given with success in gastro intestinal affections, particularly the summer diarrhoea of children.

Dose—5 to 20 grains = 0.32 to 1.3 gramme

Prescribing Notes—Given in cachets, or in a mixture suspended with Water. The salt is dissociated by contact with Water, and if an alkaline Carbonate be present, the mixture effervesces, in such cases it is better to prescribe *Bismuth Carbonate* and *Sodium Salicylate*.

Foreign Pharmacopœias—Official in Aust, Belg, Ger, Ital, Jap and Swiss, 63 p.c. of Bismuth Oxide, Dutch, 60 to 65 p.c., Fr and Mex, 61 p.c., Russ, 60 p.c., Dan and Swed 60 p.c., Norw and Span, no p.c. given, U.S., 62 to 64 p.c. Not in the others.

Tests—Bismuth Salicylate, when dissolved in diluted Hydrochloric Acid, and the Salicylic Acid separated, yields the distinctive tests for Bismuth which are mentioned under Bismuth Carbonate. It gives a violet coloration when treated with diluted Ferric Chloride Test-solution. The salt undergoes dissociation rapidly and even Alcohol (90 p.c.) causes the liberation of a certain amount of free Salicylic Acid notwithstanding the official requirement that such alcoholic liquid should not give a violet coloration with Ferric Chloride Test-solution. It is officially required to contain 98.59 p.c. of Bismuth Salicylate as gravimetrically determined by conversion into Bismuth Sulphide, 1 gramme of the Salicylate is required to yield 0.7 gramme of the Sulphide. When gravimetrically determined as Oxide it is officially required to yield 96.35 to 99.46 p.c. of Bismuth Salicylate. It will thus be seen that the amount of Bismuth Salicylate calculated from the Sulphide determination does not agree with that calculated from the Oxide determination. The U.S.P. requires it to yield not less than 62 p.c. nor more than 66 p.c. of Bismuth Oxide, when ignited as described in the small type below under the heading of Gravimetric Determination. This amount of Bismuth Oxide calculates out to not less than 96.4 p.c. nor more than 102 p.c. of Bismuth Salicylate. The P.G. requires it to leave not less than 63 p.c. of Bismuth Oxide when the salt is calcined as described in the same, corresponding to a calculated figure of not less than 97 p.c. of Bismuth Salicylate.

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead, Magnesium, Silver and Zinc, Selenium and Tellurium, Chlorides, Nitrates and Sulphates, free Salicylic Acid.

The detection of the majority of these impurities is referred to under Bismuth Carbonate, in carrying out the tests for them the Bismuth Oxide left on the ignition of the Salicylate should be dissolved in Nitric Acid, and the major portion of the Bismuth removed as an oxy-salt. With regard to the detection of free Salicylic Acid the methods adopted vary. The employment of Alcohol (90 p.c.) as recommended in the B.P. results in the liberation of a sufficient amount of free Salicylic Acid to give a pronounced coloration with Ferric Chloride Test-solution. Chloroform is a more appropriate solvent and the U.S.P. test is carried out with this menstruum, a weighed quantity of 1 gramme of the salt being shaken with 5 c.c. of Chloroform and the chloroformic solution filtered into an equal volume of Water containing 3 drops of the Ferric Chloride Test-solution, when no violet zone should form at the junction of the two liquids within one minute. The P.G. requires

that a weighed quantity of 0.5 gramme of the salt when shaken with 5 c.c. of water shall yield a filtrate which produces no reddening of blue Litmus paper, but does not also test the solution with Ferric Chloride Test-solution.

The Salicylic Acid liberated when the salt is treated with an acid, when filtered off, washed free from mineral acid and carefully dried, should possess the melting point and answer the tests given under Acidum Salicylicum and should otherwise conform to the tests of purity given for this acid. The *BP* and *PG* formulate no such requirement, the *USP* has carefully noted this.

The three Pharmacopœias differ widely in the test adopted for the detection of Nitrates. The *USP* uses a mixture of Bismuth Salicylate, Sodium Salicylate and Sulphuric Acid, the *PG* test depends upon the reduction of the Nitrates to Ammonia by the use of Zinc foil and powdered Iron and its liberation by Sodium Hydroxide Solution (15 p.c.), the *BP* depends upon the formation of Nitrous Oxide and liberation of red fumes when the salt is warmed with Sulphuric Acid and Copper, the *USP* and *PG* tests are compared in detail in small type below under the heading of Sodium Salicylate and Sulphuric Acid and Zinc foil, powdered Iron, and Sodium Hydroxide. The most delicate is that of the *USP*. Both the *USP* and *PG* employ Bettendorfs test as a means of detecting Arsenic. The Uranium Nitrate test for distinguishing it from Carbolates and Sulpho carbolates is peculiar to the *BP*, and is commented upon in the large type under Acidum Salicylicum.

Sodium Salicylate and Sulphuric Acid—0.05 gramme triturated with 0.1 gramme of Sodium Salicylate and 5 c.c. of Water, carefully poured as a layer over 5 c.c. of Sulphuric Acid (free from Nitrous Compounds) should not immediately form a pink to brownish red zone, *USP*.

Zinc Foil, Powdered Iron, and Sodium Hydroxide—On warming 0.5 gramme Bismuth Subsalicylate with 5 c.c. Sodium Hydroxide Solution and the addition of 0.5 gramme of Zinc foil and reduced Iron, Ammonia gas should not be evolved, *PG*.

Gravimetric Determination—If 1 gramme of Bismuth Subsalicylate be calcined and the residue dissolved in Nitric Acid, this solution carefully evaporated, and the residue again calcined, a final residue of at least 0.63 gramme of Bismuth Oxide should be obtained, *PG*. The *USP* directs the use of 5 c.c. of Nitric Acid in above test, adding it to the residue drop by drop until solution is complete. The final residue obtained as above should weigh not less than 0.62 gramme and not more than 0.66 gramme.

Stannous Chloride—The residue of Bismuth Oxide obtained when 2 grammes of Bismuth Subsalicylate is ignited as described above, should not respond to Bettendorfs test for Arsenic, *USP*. A mixture of 1 gramme of Bismuth Subsalicylate and 3 c.c. of Stannous Chloride T.S. should not assume a dark colour in the course of an hour, *PG*.

Not Official

BISMUTHI CERII SALICYLAS—A reddish white powder, insoluble in Water and Alcohol (90 p.c.). Recommended in diarrhoea and dysentery.

Dose—5 grains = 0.32 gramme.

The following mixture was proposed by the Royal College of Physicians for use during the prevalence of cholera in 1892—*L*, '92, p. 682—

Cholera Mixture—Bismuthi et Cerni Salicylas, 5 grains, Mist Cretæ Aromat, 1 fl oz, Tinct Camph Co, $\frac{1}{2}$ fl drim, Tinct Chloroformi Co, 20 drops, Spirit Ammon Aromat, 20 drops, Ess Menth Pip, 10 drops

Should this mixture disagree, or in 24 hours fail to give relief, the following mixture should be substituted and taken in 1 oz doses every 3 or 4 hours—

Acid Sulph Aromat, 15 drops, Tinct Camph Co, $\frac{1}{2}$ drim, Tinct Chloroformi Co, 20 drops, Tinct Coto, 20 drops, Syrupi Aulanti Flor, 1 drim, Aq. Menth Pip ad 1 oz

BISMUTHI SUBNITRAS.

BISMUTH OXYNITRATE

$\text{BiONO}_3 \cdot \text{H}_2\text{O}$, eq 302.64

FR, SOUSNITRAT DE BISMUTH, GFR, BASISCHES WISMUTHNITRAT, ITAL, SOTTO-NITRATO DI BISMUTO, SPAN, NITRATO (SUB) BISMUTICO

A heavy, white, odourless, crystalline powder, which may be prepared from Bismuth Nitrate by the action of Water

The formula calculates into 77 p.c of Oxide, but it always contains 79 to 82 p.c. If the $\text{BiONO}_3 \cdot \text{H}_2\text{O}$ exists, it is so unstable that it could certainly not be decomposed—C P '85, 561

Although Mr David Howard called attention to the inaccuracy of the formula given in B P '85, the error is repeated in B P '98. It is also at variance with the official test, which requires that it should yield 84 p.c of Bismuth Sulphide

Solubility—Insoluble in Water Insoluble in Alcohol (90 p.c). Soluble in Hydrochloric and in Nitric Acid

Medicinal Properties.—Sedative and astringent both internally and externally. It is highly useful in pyrosis, all forms of vomiting and irritative dyspepsia, in gastric ulcer, also in diarrhoea from any cause, usually combined with Soda, Magnesia, Opium, etc., it renders the fæces leaden-grey in colour. It is recommended to be injected in gonorrhœa and leucorrhœa, 60 grains to the oz of Water, the Bismuth is mixed with an equal quantity of Glycerin or suspended with Compound Tragacanth powder. The addition of Bismuth to mixtures for diarrhoea of phthisis controls it better than other ingredients alone. As an intestinal antiseptic, *see* under Bismuthi Carbonas

Externally it is sometimes used as a cosmetic, but is more or less blackened by an impure lotion, powder, or ointment in burns, eczema, and other cases when exudation and itching are present, also as an ingredient of Ferri's Snuff in acute coryza and chronic rhinitis

Has been recommended as a dressing for wounds—L '85, 11 684, T G '85, 266, B M J '01, 11 811

Dose.—5 to 20 grains = 0.32 to 1.3 gramme.

Prescribing Notes—When prescribed in a mixture, it should be suspended with Compound Powder of Tragacanth, 1 drim. in a 6-oz mixture. *See* Bismuthi Carbonas

As Bismuth Oxynitrate in Water slowly parts with its Nitric Acid, the mixture is always acid, and this somewhat interferes with its suspension, and when prepared with Sodium Bicarbonate it causes a slight but steady evolution of Carbonic

Acid, which may cause the bottle to burst, these objections do not apply to the Bismuth Carbonate, which is therefore preferable in mixtures

Incompatibles—Effervescence ensues if prescribed in Water with Alkaline Bicarbonates With Potassium Iodide double decomposition slowly ensues

Official Preparations—Used in the preparation of Liquor Bismuthi et Ammonii Citratis, and Bismuthi Oxidum

Not Official—Fenier's Snuff, Elixir Bismuthi, Glyceritum Bismuthi, Liquor Bismuthi Conc, Lotio Bismuthi, Mistura Bismuthi Comp, Mistura Bismuthi Composita cum Pepsino, Mistura Bismuthi Composita cum Morphina, Unguentum Bismuthi Oleatis, and Unguentum Bismuthi

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests—The distinguishing tests for Bismuth are given in large type under Bismuth Carbonate, and the Subnitrate when dissolved in dilute Nitric Acid should conform to these When heated in a porcelain crucible nitrous vapours are evolved and a white residue of Bismuth Oxide remains, red fumes are evolved when a little of the salt is warmed with Sulphuric Acid and metallic Copper, after separation of the Bismuth the filtrate affords when mixed with an equal volume of Sulphuric Acid, cooled and poured gently on to the surface of Ferrous Sulphate Solution a brown ring at the junction of the two fluids It is officially required to contain 99.68 p.c. of Bismuth Subnitrate, as gravimetrically determined by conversion into Bismuth Sulphide 1 gramme of the Subnitrate is required to yield 0.84 gramme, corresponding to 84 p.c. of the Sulphide The criticism of this method of determination appearing under Bismuth Carbonate applies with still greater force here, as owing to the oxidising influence of Nitric Acid, Sulphur is almost certain to be precipitated along with the Bismuth Sulphide The *USP* and *PG* employ the Oxide method of determination, the former requiring that the salt shall yield not less than 80 p.c. of pure Bismuth Oxide, the latter that it shall yield from 79 to 82 p.c. of Bismuth Oxide Bismuth Subnitrate is distinguished from the Carbonate by being soluble without effervescence in diluted Nitric Acid, from the Oxychloride by dissolving in Acetic Acid

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead, Magnesium, Silver, and Zinc, Chlorides and Sulphates, Carbonates, Selenium or Tellurium, Calcium Phosphate A standard of 2 parts per 1,000,000 is suggested (*CD '08*, i 795) for Arsenic With the exception of Carbonates and Calcium Phosphate the remarks upon the detection of these impurities appearing under Bismuth Carbonate are also applicable here The presence of Carbonates is shown by effervescence produced when the salt is dissolved in Nitric Acid, Calcium Phosphate by a precipitate or opalescence produced when a solution of 1 gramme of the salt in Nitric Acid is mixed with a solution containing twice this weight of Citric Acid and sufficient Ammonia Solution to produce a decided alkalinity Neither the *USP* nor the *PG* include a similar test The *USP* requires that no residue should be left when the salt is dissolved in warm Nitric Acid, indicating the absence of foreign salts

Diluted Sulphuric Acid—The *PG* directs that 0.5 grammes should form a clear solution at ordinary temperature with 25 c.c. of Diluted Sulphuric Acid without evolution of Carbon Dioxide, indicating the absence of Lead and Carbonates.

Stannous Chloride Solution—If 1 gramme of Bismuth Subnitrate be heated until vapours cease to be given off, the residue cooled and dissolved in a little Hydrochloric Acid, this solution, with double its volume of Stannous Chloride Solution, should not assume a dark colour in the course of an hour. *PG*. The *USP* requires that the residue, obtained by heating 2 grammes of Bismuth Subnitrate in a porcelain crucible until nitrous vapours cease to be given off, should weigh not less than 1.6 grammes and should not respond to Bettendorfs test for Arsenic.

Silver Nitrate Solution—0.5 grammes of Bismuth Subnitrate dissolved in 5 c.c. of Nitric Acid should give a clear solution which is not rendered more than opalescent by 5 c.c. of Silver Nitrate Solution, *PG*.

Barium Nitrate Solution—0.5 grammes should dissolve to a clear solution in 5 c.c. of Nitric Acid, and be unaffected by 0.5 c.c. of Barium Nitrate Solution diluted with an equal quantity of Water, *PG*.

Potassium or Sodium Hydroxide Solution—Warmed with Sodium Hydroxide Solution in excess, Bismuth Subnitrate should not evolve Ammonia, *PG*. The *USP* directs that 1 gramme of the salt be boiled with 5 c.c. Potassium Hydroxide TS.

Gravimetric Determination—The *PG* requires that 100 parts of Bismuth Subnitrate heated until the evolution of yellowish red fumes ceases, should yield from 79 to 82 parts of residue, and the *USP* that 2 grammes, heated in a porcelain crucible until nitrous vapours cease to be evolved, should yield a residue weighing when cold not less than 1.6 grammes.

Preparation

LIQUOR BISMUTHI ET AMMONII CITRATIS. SOLUTION OF BISMUTH AND AMMONIUM CITRATE *BP Syn*—LIQUOR BISMUTHI.

A clear, colourless liquid possessing generally a faint odour of Ammonia and a slight metallic taste, 1 fl. drm is equal to rather less than 3 grains of Bismuth Oxide.

Dose— $\frac{1}{2}$ to 1 fl. drm = 1.8 to 3.6 c.c.

A formula is given in *USNF*, using Glycerite of Bismuth (see below), Alcohol and Distilled Water. 1 fl. drm equals 1 gram Bismuth and Ammonium Citrate.

Tests.—Bismuth and Ammonium Citrate Solution should possess a specific gravity of 1.070. It should be faintly alkaline in reaction towards red Litmus paper, and when warmed with an excess of Potassium or Sodium Hydroxide Solution should yield a strong ammoniacal odour and a white precipitate. It is required to contain 9 p.c. by weight equivalent to 9.61 p.c. w/v of Bismuth Ammonium Citrate $\text{BiC}_2\text{H}_3\text{O}_7(\text{NH}_3)_2$, eq. 445.64, as gravimetrically determined by precipitation of the Bismuth as Sulphide. The amount of Bismuth Sulphide yielded by the *BP* test is 5.14 p.c. by weight equivalent to 5.5 p.c. w/v, the corresponding amount of Bismuth Oxide being 4.65 p.c. by weight equivalent to 5 p.c. w/v.

The more general occurring impurities are Arsenic, Copper, Iron, Lead, Silver, Selenium or Tellurium. Of these the more important are Arsenic and Lead, Selenium and Tellurium. To detect these

impurities the liquor is evaporated to dryness, ignited and the residue redissolved in diluted Nitric Acid. The presence of Arsenic may be detected by Bettendorf's test, Lead by diluted Sulphuric Acid after separation of the Bismuth as an oxy-salt, and Selenium or Tellurium by the Sodium Hypophosphite or Sodium Sulphite tests

A good deal of controversy has ranged round the official method of preparation. The earlier criticisms (*C D* '98, 1 620) were apparently favourable to the new formula. Later (*C D* '98, 1 955) it has been pointed out that when the process is carried out strictly according to the *B P* directions that considerable difficulty arises in taking into solution the whole of the Bismuth Citrate precipitate, it being shown that the quantity of Ammonium Citrate ordered is the main factor in determining this insolubility. An increase in the amount of Ammonium Citrate is also suggested (*C D* '99, 11 211, 233, *P J* '99, 11 101, 116), the addition of *Liquor Ammonii Citratis* (after the solution of the precipitate), in the proportion of 8 fl. oz (or 400 c c) and then dilution to 20 fl. oz (or 1000 c c) is recommended. Amongst other modifications suggested in the same reference are the omission of the addition of Distilled Water until the liquid is very faintly opalescent, the use of 236 grains instead of 175 grains of Potassium Carbonate per pint of liquor, the latter quantity being inadequate to neutralise the free Nitric Acid and a corresponding amount of Bismuth being lost in the acid liquid, the non-dilution of the Bismuth Oxynitrate and Nitric Acid mixture, the reversion of the order of mixing, the Bismuth mixture being added to the Potassium Citrate and Carbonate instead of *vice versa*, and the use of a definite quantity of Water for dissolving the Potassium salts. The precipitate thus produced is easily washed, and is perfectly soluble. Samples containing the exact amount of Ammonium Citrate recommended by the *B P* formula deposit when dispensed with alkali Bicarbonates, solutions containing an additional quantity remain clear.

The conclusion drawn (*C D* '02, 1 852, '02, 11 312, *P J*, '02, 11 135) is that Bismuth Citrate is an acid and not a salt, being Citric Acid with one of the Hydrogen atoms replaced by Bismuthyl. Its acid properties are shown by it forming salts with Ammonia and yielding an effervescence with alkali Carbonates and Bicarbonates. Determinations of the combining weight of the acid are also recorded in support of this theory. The following process is suggested:—629 grains of Bismuth Subnitrate mixed in a mortar with $1\frac{1}{2}$ fl. oz of Water, is set aside for 2 hours, occasionally stirring, or until the mixture yields a clear solution with Ammonia Solution, sufficient of the latter is then added to form a clear solution and the mixture diluted to 20 fl. oz with water and filtered. An alteration in the official title to *Liquor Ammonii Bismuthyl-Citratis* is recommended. The chief drawback to the official method of preparation appears (*P J* '99, 11 604) to be the use of a smaller quantity of Potassium Citrate than is necessary for converting the whole of the Bismuth into Citrate. The method recommended in this reference is to dissolve 70 grammes Bismuth Oxynitrate in diluted Nitric Acid by gently warming, to add 50 grammes of Citric Acid dissolved in a little water (and if a

Carbonate is used two-thirds of it may be mixed with the Citric Acid), and a solution of 103 grammes of Potassium Bicarbonate or 86 grammes of Sodium Bicarbonate and dilute with hot Water to 1000 c c, cool, filter and wash free from Nitrate. The precipitate is dissolved in 60 c c of Liquor Ammonia *B P* diluted to 200 c c with Water and made up to 1000 c c or to a specific gravity of 1.070.

Not Official.

ELIXIR BISMUTHI—Bismuth and Ammonium Citrate, 3.50, Distilled Water, hot, 6, Water of Ammonia (*U S P*), *q s*, Aromatic Elixir, *q s* to produce 100—*U S N F* 1896.

This has been incorporated in the *B P C*

Glycerite of Bismuth (*N F*), 12.5, Glycerin, 12.5, Water, 25, Aromatic Elixir (*U S P*), 50—*U S N F* 1906.

BISMUTH ET AMMONII CITRAS EFFERVESCENS 2 grains are contained in 1 dm.

FERRI'S SNUFF—Bismuth Subnitrate, 6 dm, Morphine Hydrochloride, 2 grains, Gum Acacia, in powder, 2 dm—*L*, '76, 1525.

It is described as a speedy and efficacious remedy for a recent cold in the head, each time the nostrils are cleared another pinch should be taken, using it frequently at first. One quarter to one half of this formula may be used in the twenty-four hours.

Glass insufflators are made to blow it up the nostrils.

This has been incorporated in the *B P C* as follows—

Insufflatio Bismuthi et Morphinae—Bismuth Subnitrate, 75, Morphine Hydrochloride 0.40, Gum Acacia, in powder, to make 100.

GLYCERITUM BISMUTHI—A solution of Bismuth and Ammonium Citrate in Ammonia and Water containing Glycerin. 1 fl dm contains 16 grains of the salt—*U S N F* 1896.

4 c c (1 fl dm) contains about 1 gm (16 grains) of Bismuth and Sodium Tartrate—*U S N F* 1906.

LIQUOR BISMUTHI CONC—Dissolve 7 of Bismuth Subnitrate in 10 of equal volumes of Nitric Acid and Distilled Water with a gentle heat, when cold add first a solution of Citric Acid 5 in Distilled Water 7, and subsequently stirring in a solution of Sodium Bicarbonate 8½ in Distilled Water 7. Wash the precipitate free from Nitrates, and after draining dissolve it in solution of Ammonia 6, or a sufficient, and add solution of Ammonium Citrate 12, and Distilled Water, *q s* to yield 50—*B P C Formulary* 1901.

Incorporated in the *B P C*

LOTIO BISMUTHI—Bismuth Subnitrate, 10 grains, Water, 1 fl oz. A sedative lotion in cases of eczema.

MISTURA BISMUTHI COMPOSITA—Compound—Caradomoms, 9 fl oz, Chloroform, 70 minims, Liquid Extract of Nux Vomica, 85 minims, Diluted Hydrocyanic Acid, 320 minims. Mix and add Concentrated Solution of Bismuth, 15 fl oz, Morphine Hydrochloride, 8 grains dissolved in 4 fl dm of Distilled Water, add finally Distilled Water, *q s* to yield 20 fl oz.

Each fl dm contains 2 minims of Diluted Hydrocyanic Acid, ½ grain of Morphine Hydrochloride, and the equivalent of 5 minims of Tincture of Nux Vomica—*B P C Formulary* 1901.

Dose—20 to 30 minims=1.2 to 1.8 c c

Mistura Bismuthi Composita—Bismuth Citrate, 320 grains; Solution of Ammonia, *q s*, Chloroform, 32 minims, Tincture of Nux Vomica 1 fl oz, Diluted Hydrocyanic Acid 128 minims, Solution of Carmine (Martindale), 32 minims, Distilled Water, *q s* to produce 8 fl oz. Rub the Bismuth Citrate with a little Water, add Solution of Ammonia until salt is just dissolved

and make up to 6 oz with Distilled Water. Dissolve the Chloroform in the Tincture of Nux Vomica and add to the Bismuth Solution, then add the Solution of Carmine and filter, wash the filter paper with sufficient Distilled Water to produce with the Hydrocyanic Acid 8 fl oz of finished product. Each fl drim is equivalent to 1 drim of the *B P* Bismuth Solution, 10 minims of Spirit of Chloroform, $7\frac{1}{2}$ minims of Tincture of Nux Vomica, and 2 minims of Diluted Hydrocyanic Acid. Dose — $\frac{1}{2}$ to 1 drim — *Bournemouth Formulary*

This has been incorporated in the *B P C*

MISTURA BISMUTHI COMPOSITA CUM PEPSINO — Bismuth Citrate, 320 grains, Solution of Ammonia, *q s*, Soluble Scale Pepsin, 64 grains; Chloroform, 32 minims, Tincture of Nux Vomica, 1 fl oz, Diluted Hydrocyanic Acid, 128 minims, Solution of Carmine (Martindale), 32 minims, Distilled Water, *q s* to produce 8 fl oz. Rub the Bismuth Citrate with a little Water, add Solution of Ammonia until salt is just dissolved, and make up to 4 oz with Distilled Water. Dissolve the Pepsin in 2 oz of Water and add to the Bismuth Solution, then add the Chloroform dissolved in the Tincture of Nux Vomica, and the Carmine Solution, filter, and wash the filter paper with sufficient Water to produce with the Hydrocyanic Acid 8 fl oz of finished product. Each fl drim is equivalent to Solution of Bismuth, 1 drim, Spirit of Chloroform, 16 minims, Tincture of Nux Vomica, $7\frac{1}{2}$ minims, Pepsin, 1 grain, Hydrocyanic Acid, 2 minims — *Bournemouth Formulary*

This has been incorporated in the *B P C*

MISTURA BISMUTHI COMPOSITA CUM MORPHINA — Morphine Hydrochloride, 1 grain, Compound Bismuth Mixture, 3 fl oz. Each fl drim contains $\frac{1}{24}$ grain of Morphine Hydrochloride. Dose — $\frac{1}{2}$ to 1 drim — *Bournemouth Formulary*

This has been incorporated in the *B P C*

UNGUENTUM BISMUTHI — Bismuth Subnitrate, 60 grains, Lard, 1 oz. Bismuth Subnitrate, 12 5, Lard, 87 5 — *B P C*

BISMUTHI BENZOAS — A white powder, without taste, almost insoluble in Water. Given internally as an antiseptic and sedative. Used externally as an antiseptic dusting powder.

Dose — 5 to 20 grains = 0 32 to 1 3 gramme

Foreign Pharmacopœias — Official in Mex, not in the others

Tests — When dissolved in diluted Hydrochloric Acid and separated from the precipitated Benzoic Acid, the filtrate should answer the tests distinctive of Bismuth given under Bismuth Carbonate. When shaken with a few drops of Ferric Chloride Test-solution a buff coloration is produced. The Benzoic Acid separated from the salt should possess the melting point and conform to the tests for identity and purity given under Benzoic Acid. The salt should leave from 60 to 70 p c of Bismuth Oxide after ignition at a low red heat.

It should be free from the impurities mentioned under Bismuth Carbonate

BISMUTH CITRAS — A white amorphous, odourless and tasteless powder

Solubility — Insoluble in Water, readily in Solution of Ammonia, and in solutions of alkali citrates

Medicinal Properties — Similar to the Subnitrate

Dose — 2 to 5 grains = 0 13 to 0 32 gramme

Foreign Pharmacopœias — Official in U S. Not in the others

Tests — The residue obtained on ignition, when dissolved in warm Nitric Acid yields the tests distinctive of Bismuth given under Bismuth Carbonate. The salt chars when first heated and the residue left on ignition is more or less black in colour with a yellow surface. If to the solution of Bismuth Citrate in Ammonia Solution sufficient Hydrogen Sulphide be added to precipitate the whole of the Bismuth, the filtrate, when freed from Hydrogen Sulphide and boiled with an excess of Calcium Hydroxide Solution yields a white precipitate. The salt is official in the *U S P* and is required to contain not less than $\frac{1}{4}$ 75 p c

nor more than 99.17 p.c. of pure Bismuth Citrate, as gravimetrically determined by weighing.

As regards arsenic, it should be free from the impurities mentioned under Bismuth Carbonate and when the Bismuth is separated by Hydrogen Sulphide, the latter removed, the filtrate when mixed with an equal volume of concentrated Sulphuric Acid, and cooled should produce no brown or brownish-black colour round a crystal of Ferrous Sulphate.

BISMUTHI ET AMMONII CITRAS—Small shining translucent scales, which yield Ammonia when warmed with a solution of a fixed alkali, and gradually lose Ammonia on exposure to the air.

Solubility—1 in 1 of Water, sparingly in Alcohol (90 p.c.)

Dose.—2 to 5 grains = 0.13 to 0.32 gramme

Foreign Pharmacopœias.—Official in U.S. Not in the others.

Tests—A blackish residue with a yellow surface remains when the salt is ignited, and a solution of this residue in warm Nitric Acid yields the tests distinctive of Bismuth given under Bismuth Carbonate. When heated with an excess of Potassium or Sodium Hydroxide Solution, the salt evolves a strong odour of Ammonia and yields a white precipitate when the Bismuth is separated from its solution by means of.

and the excess of the latter is removed by heat, the filtrate when boiled with an excess of Calcium Hydroxide Solution yields a white precipitate. The U.S.P. requires the salt to yield not less than 46 p.c. nor more than 50 p.c. of pure Bismuth Oxide as gravimetrically determined by ignition and oxidation with Nitric acid.

It should be free from the impurities mentioned under Bismuth Carbonate, should give a negative test for Arsenic, and when separated from the Bismuth and from the excess of the latter by boiling should, when mixed with an equal volume of concentrated Sulphuric Acid and cooled, yield no brown or brownish-black coloration to a crystal of Ferrous Sulphate dropped into the mixture.

BISMUTHI NITRAS ($\text{Bi}(\text{NO}_3)_3 \cdot 5 \text{H}_2\text{O}$, eq. 481.44)—In colourless transparent crystals. Decomposed by Water, giving a white precipitate of Subnitrate. Soluble in Glycerin, but is slowly deposited from the solution when Water is added.

A glycerole can be made containing 60 grains to the oz.; but as an application in skin diseases the strength should in most cases not exceed 10 grains to the oz.—*M.T.* 76 ii 646.

The salt should be dissolved without the application of heat.

Official in Fr. (Azotate Neutre de Bismuth)

BISMUTHI OLEAS—Crystallised Bismuth Nitrate, 280 grains, dissolve cold in Glycerin 4 oz. by weight, add slowly Solution of Sodium Oleate, 20 fl. oz.; warm gently, wash by decantation, collect and dry. It forms a pearly-grey soft bland substance.

Medicinal Properties. It is a reliable application in pustular eruptions and hyperæmia of the skin.—*B.M.J.* '84, ii 751.

Unguentum Bismuthi Oleatis (Sir T. McCall Anderson)—Oxide of Bismuth, 1 drim, Oleic Acid, 1 drim, White Wax, 1½ drim, Vaseline, 9 drim—*Pharm. Form.*

Bismuth Oleate, 10, Soft Paraffin, 90.—*B.P.C.*

BISMUTH-PHENOL (Bismuth Phenate)—Prepared by adding a solution of Phenol in an alkali, to a solution of Bismuth Oxynitrate. A greyish-brown amorphous powder, insoluble in Water and Alcohol (90 p.c.). Recommended as an intestinal antiseptic.—*J.J.* (3) xxiv 182, *C.D.* 93, ii 576.

Dose.—5 to 20 grains = 0.32 to 1.3 gramme.

BISMUTHI SUBGALLAS (Dermatol)—A light yellow amorphous insoluble powder, introduced as an odourless substitute for Iodoform.

Sometimes causes symptoms of Bismuth poisoning.

Given for gastric ulcer and diarrhoea in doses of 8 to 30 grains twice a day —
L '97, 11 404

Solubility—Insoluble in Water, Alcohol (90 p c) and in Ether Soluble with decomposition in mineral acids, and readily soluble in alkali Hydroxide Solutions

Foreign Pharmacopœias—Official in Austr, 58 to 55 p c of Bi_2O_3 , Belg, 52 p c, Dutch, 52 to 56 p c, Dan, Ger, Swed and Swiss, 52 p c, Fr, 56 45 p c, Ital, not less than 55 p c, Jap, 51 p c, Mex and Russ, no p c given, Russ also includes a Tannate, Span, 56 66 p c, U S, 52 to 57 p c Not in the others

Tests—When strongly heated it chars, and on ignition leaves a yellow residue, which dissolves in Hydrochloric and in Nitric Acids, yielding a solution which answers the tests distinctive of Bismuth given under Bismuth Carbonate After complete separation of the Bismuth by means of Hydrogen Sulphide and the removal of the latter by boiling, the cold filtrate yields with a drop of Ferric Chloride Test solution a bluish-black coloration The salt is official in the *U S P* and in the *P G*, the former requires it to yield not less than 52 p c nor more than 57 p c of pure Bismuth Oxide, the latter not less than 52 p c of Bismuth Oxide

The more generally occurring impurities are Arsenic and those usually associated with Bismuth, Ammonium salts, Carbonate and Nitrate, free Gallic Acid Bettendorf's test is employed in both Pharmacopœias as a test for Arsenic The methods of detecting the impurities usually associated with Bismuth are given under Bismuth Carbonate The *U S P* and *P G* both use Alcohol as a solvent for free Gallic Acid, the former being content with the neutrality of the alcoholic solution towards blue Litmus paper as ensuring its absence, the latter evaporating the alcoholic solution to dryness, when no weighable residue should remain The *U S P* employs the Sulphuric Acid and Ferrous Sulphate test for Nitrate, the *P G* the reduction with Zinc and powdered Iron, when on boiling with Sodium Hydroxide Solution (15 p c), no evolution of Ammonia should take place

Bismal (Bismuth Methylendigallate)—Introduced as an astringent for internal administration in cases of diarrhoea Insoluble in Water

Dose—1 to 4 grains = 0 06 to 0 26 gramme

BISMUTHI OXYIODOGALLAS (Aiol, Airoform and Airogen)—A bulky greyish powder, odourless and tasteless, insoluble in Water and Alcohol, soluble in dilute mineral acids and in Sodium or Potassium Hydroxide Solution It is gradually converted into a more basic salt by the action of light and moist air, and should therefore be kept in well-stoppered dark amber-tinted glass bottles A combination of Dermatol with Iodine, introduced as a substitute for Iodoform, has attracted a good deal of attention as an antiseptic dressing Used as a dusting powder for ulcers, also mixed with Vaseline or anhydrous Lanolin —*B M J* '98, 1 144, *L* '99, 1 240

Sometimes badly tolerated —*B M J E* '97, 11 43

Comparative experiments with Aiol, Dermatol and Iodoform —*B M J E* '97, 1 67

Foreign Pharmacopœias—Official in Belg and Swiss Not in the others

Tests—A solution in dilute Hydrochloric Acid gives with Hydrogen Sulphide a black precipitate, and if this precipitate be separated, washed and dissolved in Nitric Acid the solutions should yield the tests distinctive of Bismuth given under Bismuth Carbonate A small quantity of the salt warmed with a few drops of concentrated Sulphuric Acid evolves violet vapours of Iodine, its Hydrochloric Acid Solution when treated with Chlorine Water and shaken with Carbon Bisulphide, colours the latter violet A solution in very dilute Hydrochloric Acid gives with Ferric Chloride Test-solution a dark green coloration It should contain about 46 p c by weight of Bismuth Oxide, and about 24 0 p c by weight of Iodine

It should be free from the impurities mentioned under Bismuth Carbonate

BISMUTHI BETA-NAPHTHOLAS (Oiphol) — A reddish-brown powder, insoluble in Water. Recommended as an intestinal antiseptic and astringent, both for adults and children.

Dose — 5 to 90 grains = 0.32 to 1.3 gramme

Experiments with Bismuth Subnitrate and Beta-naphthol as intestinal antiseptics — *B M J* '95, ii 1483

BISMUTHI SUBIODIDUM — A brick-red amorphous powder, insoluble in Water.

Has been recommended as a substitute for Iodoform in the treatment of chancres and foul ulcers — *T G* '87, 612, *Y B P* '87, 286, *B M J* '89, i 783

Dose — 5 to 10 grains = 0.32 to 0.65 gramme

BISMUTHI TRIBROMOPHENOLAS (Xeroform) — A yellow powder, insoluble in Water and in Alcohol (90 p c). It has been recommended as a non-irritating antiseptic.

Used in wound dressing in the Cuban war — *L* '99, i 1509, and '99, ii 1459, *B M J E* '99, ii 88

Foreign Pharmacopœias — Official in Jap, Span and Swiss. Not in the others.

Tests — A white curdy precipitate is thrown down when a solution of the powder in Potassium Hydroxide Solution is acidified with diluted Sulphuric Acid, and if this precipitate be separated, washed and dried it should possess a melting point of 95° C (203° F). The filtrate from this precipitate should yield the tests distinctive of Bismuth given under Bismuth Carbonate. The powder should yield on ignition from 57 to 61 p. c. of Bismuth Oxide. It should be free from the more generally occurring impurities mentioned under Bismuth Carbonate and the residue remaining after ignition at a dull red heat when moistened with Nitric Acid, again ignited and when cool dissolved in Hydrochloric Acid should yield no reaction for Arsenic when tested by Bettendorf's test.

Bismone (Colloidal Bismuth Oxide), **Bismuthi Iodo-Resorcin-Sulphonas** (Anusol), supplied in Suppository form, **Bismuthi Quinolini Sulphocyanidum** (Crurin), **Bismuthi Cinchonidinæ Iodidum** (Lancet), **Bismuthi Di-thio-Salicylas** (Lithorin), **Bismuthi Oxylchloridum**, **Bismuthi Phosphas**, and **Bismuthi Sulphas**, are combinations of Bismuth, mostly insoluble in Water, which have received notice in Medical Literature.

Not Official.

BOLDO.

The Leaves and young Twigs of the *Peumus Boldus*, Mol., a native of Chili. The activity is due to a Glucoside, Boldine, and a volatile Oil (sp gr 0.918).

Foreign Pharmacopœias — Official in Mex and Span. Not in the others.

Medicinal Properties — Has been used as a liver stimulant and as a diuretic, as a stimulant to digestion, also as a hypnotic — *B M J* '85, ii 1134, '88, i 918, gastric stimulant and sedative, antispasmodic, cholagogue — *B M J E* '07, ii 72

Boldine has been given as a hypnotic in capsules containing 3 grains.

TINCTURA BOLDO. — Boldo Leaves, 1, Alcohol (60 p c), 10

Macerate seven days and filter

Dose. — 10 to 40 minims = 0.6 to 2.4 c.c.

This has been incorporated in the *B P C*.

Foreign Pharmacopœias. — Mex, 1 and 5; by weight. Not in the others.

BONE MARROW See **MEDULLA RUBRA**

BORAX.**BORAX***B P Syn*—BIBORATE OF SODIUMFR, BORATE DE SOUDE, GER, NATRIUMBORAT, ITAL, BORATO DI SODIO,
SPAN, BORATO SODICO $\text{Na}_2\text{B}_4\text{O}_{10}$, $10\text{H}_2\text{O}$, eq 379 12

Transparent colourless monoclinic prisms, usually efflorescent

Though this salt is acid in constitution, it gives alkaline reactions with Litmus and Methyl Orange Solutions

Solubility—1 in 25 of Water, 2 in 1 of boiling Water, 2 oz of Borax are dissolved by 2 fl oz of Glycerin, and the solution measures only $3\frac{1}{4}$ fl oz By the aid of 1 of Glycerin, 1 part of Borax will dissolve in 12 of Water Insoluble in Alcohol (90 p c)

Borax is decomposed by Glycerin, forming a solution which reddens Litmus paper, and effervesces with Sodium Bicarbonate

Medicinal Properties—Antiseptic and parasiticide, mildly astringent A local sedative to inflamed mucous membrane As a lotion 10 grains to the oz, as a gargle (saturated solution) about 20 grains to the oz, and as an injection in leucorrhœa and gonorrhœa The Glycerin of Borax is used as a paint for the throat, for cracked nipples, and for erythematous skin eruptions The Glycerin or Mel is used in aphthous ulceration of the tongue or buccal mucous membrane, and for mercurial salivationInternally in epilepsy (*L* '93, 11 1586, '95, 11 755), but is inferior to Bromide and has many inconveniences—*B M J E* '95, 1 4Has been recommended by some authorities in epilepsy, but (*L* '05, 1 710) unless in combination with Bromides it has not been found of much use, although a combination of Borax and Digitalis has been found serviceable in some cases of minor epilepsyThe effects of Borax on infants—*L* '07, 11 369**Dose.**—5 to 20 grains = 0 32 to 1 3 gramme**Prescribing Notes**—*For internal use it is generally given in solution Should not be prescribed with salts of Cocaine or other alkaloids***Incompatibles.**—Mineral Acids and most of their metallic salts, also alkaloidal salts Mucilage of Gum Acacia**Official Preparations**—Glycerinum Boracis and Mel Boracis**Not Official**—Gargarisma Boracis, Liquor Boracis, Liquor Sodii Boratis Compositus, Lotic Boracis, Nebula Antiseptica Alkalina, Seiler's Antiseptic, Tinctura Myrrhæ et Boracis, Trochisci Boracis, and Unguentum Boracis**Foreign Pharmacopœias**—Official in Austr, Belg, Dan, Dutch, Fr, Ger., Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S**Tests.**—The distinguishing tests for Sodium Borate are the brown coloration (changing to bluish-black on treatment with alkalis) which its acidified aqueous solution produces with Turmeric paper, the intense yellow coloration which the salt imparts to a non-luminous flame; the white scaly crystalline precipitate thrown down when a hot saturated solution of the salt is acidulated with a mineral acid, and the green coloration imparted to a non-luminous flame when the solution of this precipitate in Alcohol (90 p c) is ignited *U S P*

BISMUTHI BETA-NAPHTHOLAS (Orphol) —A reddish-brown powder, insoluble in Water. Recommended as an intestinal antiseptic and astringent, both for adults and children

Dose —5 to 20 grains = 0.32 to 1.3 gramme

Experiments with Bismuth Subnitrate and Beta-naphthol as intestinal antiseptics —*B M J* '95, n 1483

BISMUTHI SUBIODIDUM —A brick-red amorphous powder, insoluble in Water

Has been recommended as a substitute for Iodoform in the treatment of chancre and foul ulcers —*T G* '87, 612, *Y B P* '87, 286, *B M J* '89, 1 783

Dose —5 to 10 grains = 0.32 to 0.65 gramme

BISMUTHI TRIBROMOPHENOLAS (Xeroform) —A white, crystalline, insoluble in Water and in Alcohol (90 p c). It has been recommended as an irritating antiseptic.

Used in wound dressing in the Cuban war —*L* '99, 1 1509, and '99, n 1459, *B M J E* '99, 1 88

Foreign Pharmacopœias —Official in Jap, Span and Swiss. Not in the others

Tests.—A white precipitate is thrown down when a solution of the powder in Potassium Iodide solution is acidified with diluted Sulphuric Acid, and if this precipitate be separated, washed and dried it should possess a melting point of 95° C (203° F). The filtrate from this precipitate should yield the tests distinctive of Bismuth given under Bismuth Carbonate. The powder should yield on ignition from 57 to 61 p c of Bismuth Oxide. It should be free from the more generally occurring impurities mentioned under Bismuth Carbonate and the residue remaining after ignition should not become moistened with Nitric Acid, again ignited and dissolved in Hydrochloric Acid should yield no reaction for Arsenic.

Bismone (Colloidal Bismuth Oxide), **Bismuthi Iodo-Resorcin-Sulphonas** (Anasol), supplied in Suppository for **Bismuthi Quinolini Sulphocyanidum** (Cruin), **Bismuthi Cinchonidinæ Iodidum** (Cruin), **Bismuthi Di-thio-Salicylas** (Thioform), **Bismuthi Oxochloridum**, **Bismuthi Phosphas**, and **Bismuthi Sulphas**, are Water, which have received notice in Medical Literature

Not Official.

BOLDO.

The Leaves and young Twigs of the *Peumus Boldus*, Mol., a native of Chili. The activity is due to a Glucoside, Boldine, and a volatile Oil (sp. gr. 0.918)

Foreign Pharmacopœias —Official in Mex and Span. Not in the others

Medicinal Properties —Has been used as a liver stimulant and as a diuretic, as a stimulant to digestion, also as a hypnotic —*B M J* '85, n 1134, 88, 1 918, gastric stimulant and sedative, antispasmodic, cholagogue —*B M J E* '07, 1 72

Boldine has been given as a hypnotic in capsules containing 3 grains.

TINCTURA BOLDO —Boldo Leaves, 1, Alcohol (60 p c), 10

Macerate seven days and filter

Dose.—10 to 40 minims = 0.6 to 2.4 c c

This has been incorporated in the *B P C*

Foreign Pharmacopœias.—Mex, 1 and 5, by weight. Not in the others.

BONE MARROW See **MEDULLA EUBRA**,

BORAX.**BORAX***B P Syn*—BIBORATE OF SODIUM

FR, BORATE DE SOUDE, GER, NATRIUMBORAT; ITAL, BORATO DI SODIO, SPAN, BORATO SODICO

 $\text{Na}_2\text{B}_4\text{O}_7, 10\text{H}_2\text{O}$, eq 379 12

Transparent colourless monoclinic prisms, usually efflorescent

Though this salt is acid in constitution, it gives alkaline reactions with Litmus and Methyl Orange Solutions

Solubility—1 in 25 of Water, 2 in 1 of boiling Water, 2 oz of Borax are dissolved by 2 fl oz of Glycerin, and the solution measures only $3\frac{1}{4}$ fl oz By the aid of 1 of Glycerin, 1 part of Borax will dissolve in 12 of Water Insoluble in Alcohol (90 p c)

Borax is decomposed by Glycerin, forming a solution which reddens Litmus paper, and effervesces with Sodium Bicarbonate

Medicinal Properties—Antiseptic and parasiticide, mildly astringent A local sedative to inflamed mucous membrane As a lotion 10 grains to the oz, as a gargle (saturated solution) about 20 grains to the oz, and as an injection in leucorrhœa and gonorrhœa The Glycerin of Borax is used as a paint for the throat, for cracked nipples, and for erythematous skin eruptions The Glycerin or Mel is used in aphthous ulceration of the tongue or buccal mucous membrane, and for mercurial salivationInternally in epilepsy (*L* '93, 11 1586, '95, 11 755), but is inferior to Bromide and has many inconveniences—*B M J E* '95, 1 4Has been recommended by some authorities in epilepsy, but (*L* '05, 1 710) unless in combination with Bromides it has not been found of much use, although a combination of Borax and Digitalis has been found serviceable in some cases of minor epilepsyThe effects of Borax on infants—*L* '07, 11 369**Dose.**—5 to 20 grains = 0.32 to 1.3 gramme**Prescribing Notes**—*For internal use it is generally given in solution Should not be prescribed with salts of Cocaine or other alkaloids***Incompatibles**—Mineral Acids and most of their metallic salts, also alkaloidal salts Mucilage of Gum Acacia**Official Preparations**—Glycerinum Boracis and Mel Boracis**Not Official**—Gargarisma Boracis, Liquor Boracis, Liquor Sodii Boratis Compositus, Lotio Boracis, Nebula Antiseptica Alkalina, Seiler's Antiseptic, Tinctura Myrrhæ et Boracis, Trochisci Boracis, and Unguentum Boracis**Foreign Pharmacopœias**—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U.S.**Tests**—The distinguishing tests for Sodium Borate are the brown coloration (changing to bluish-black on treatment with alkalis) which its acidified aqueous solution produces with Turmeric paper, the intense yellow coloration which the salt imparts to a non-luminous flame, the white scaly crystalline precipitate thrown down when a hot saturated solution of the salt is acidulated with a mineral acid, and the green coloration imparted to a non-luminous flame when the solution of this precipitate in Alcohol (90 p c) is ignited. **U S P**

states that an aqueous solution (1 in 20) after being acidulated with Hydrochloric Acid colours blue Litmus red, yellow Turmeric paper remains unchanged until after drying

It is officially required to contain 98.57 p.c. of Sodium Pyroborate, as volumetrically determined by titration with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as the indicator of neutrality, 1 gramme neutralises 5.2 c.c. of the Volumetric Solution. Phenolphthalein Solution is of no use for this titration, and even Litmus Solution gives a rather indefinite end reaction. Although Borax is constitutionally an acid salt, Boric Acid has so little action upon the usual indicators that the Sodium Oxide can be estimated by standard acid just as if no Boric Acid were present. It has been pointed out (*PJ* '02, 1 345) that inasmuch as the salt is likely to contain Sodium Carbonate, the official process is apt to give erroneous figures. The direct determination of the Boric Acid by a double titration has been suggested. The same number of c.c. of Normal Volumetric Sodium Hydroxide Solution should be required to neutralise the Boric Acid as are required of Semi-normal Volumetric Sulphuric Acid to liberate it, the titration of the free Boric Acid being conducted in 50 p.c. Glycerin Solution. The proportions indicated in the test recommended are —1 gramme of Borax dissolved in 40 c.c. of Water should require for exact neutralisation 10.55 c.c. of Semi-normal Volumetric Sulphuric Acid and 10.55 c.c. of Methyl Orange Solution as an indicator of neutrality, after boiling and adding 50 grammes of Glycerin, 10.55 c.c. of Normal Sodium Hydroxide Solution should be required to exactly neutralise, Phenolphthalein Solution being employed as the indicator of neutrality. Neither *PG* nor *USP* gives any quantitative test for Borax.

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead, Magnesium, Bicarbonates, Carbonates and Nitrates, Phosphates, Chlorides and Sulphates. The *BP* groups these collectively under the usual elastic expression 'A standard of 5 parts per 1,000,000 is suggested' (*CD* '08, 1 726) for Arsenic. Tests for heavy metals, Carbonate and Bicarbonate, Nitrate and Phosphate appear in the *USP*, but no tests for Calcium, Magnesium, or Sulphate. The *PG* includes, in addition to tests for heavy metals, specific tests for Calcium, Iron, Chlorides and Sulphates. A 2 p.c. aqueous solution of the salt when acidified with Hydrochloric Acid should be unaffected by Hydrogen Sulphide Solution, indicating the absence of Arsenic, Copper and Lead. 50 c.c. of a 1 in 50 aqueous solution, after the addition of a few drops of Hydrochloric Acid should not yield an immediate coloration on the addition of 0.5 c.c. of Potassium Ferrocyanide Solution, indicating the absence of more than a faint trace of Iron. The 1 in 50 aqueous solution acidified with Acetic Acid should yield no turbidity on the addition of Ammonium Oxalate Solution, indicating the absence of Calcium, and when allowed to stand some time, this solution is filtered, should not yield on the addition of Ammonium Phosphate Solution any turbidity or precipitate, indicating the absence of Magnesium. The aqueous solution should not effervesce on the addition of

mineral acid, indicating the absence of Bicarbonates and Carbonates A 1 in 50 aqueous solution should not be rendered turbid on the addition of either Silver Nitrate or Barium Nitrate Solution, indicating the absence of Chlorides and Sulphates, nor should it be rendered turbid by Magnesium Ammonio-sulphate Solution, indicating the absence of Phosphates The *U S P* employs the Indigo test for Nitrates, requiring that if 1 gramme be dissolved in 20 c c of diluted Sulphuric Acid, by the aid of heat, and 3 drops of Indigo Test-solution be added, the blue coloration should not be discharged after heating for 10 minutes on a water-bath

Preparations

GLYCERINUM BORACIS GLYCERIN OF BORAX

Borax, 1, Glycerin, 6 (by weight 1 in $8\frac{1}{2}$, measure 1 in $6\frac{1}{2}$)

This is not merely a solution of Borax in Glycerin, the Glycerin splits up the Biorate into free Boric Acid and a more basic Borate with secondary reactions. It reddens blue Litmus paper, and effervesces on the addition of Sodium Bicarbonate

Dose — $\frac{1}{2}$ to $1\frac{1}{2}$ fl drm = 1 8 to 5 4 c c

20 minims given in diarrhoea of infants — *L* '89, 11 739

Foreign Pharmacopœias — Official in Mex (Glycerina Boratada), 1 and 19, Dan and Norw (*Linctus boracinus*), 1 and 9, all by weight Not in the others

MEL BORACIS BORAX HONEY

Borax, 2, Glycerin (by weight), 1, Clarified Honey (by weight), 16
(about 1 in 7 by volume)

Foreign Pharmacopœias — Official in Austr, 1 in 20, Mex (*Colutorio boratado*), Borax 1, Honey 1, Swiss, 1 in 10, the ingredients vary slightly Not in the others

Not Official.

GARGARISMA BORACIS — Borax, 1, Water, to 20 — *St Thomas's*

Borax, 4, Distilled Water, to produce 100 — *B P C*

Borax, 10 grains, "Glycerin, 30 minims, Distilled Water, to 1 fl oz — *St George's*

LIQUOR BORACIS (Thompson's Fluid) — Borax, 1, Glycerin, 2, Water, 2 $\frac{1}{2}$ oz to be mixed with 4 fl oz of warm Water before use — *Guy's*

LIQUOR SODII BORATIS COMPOSITUS (Dobell's Solution) (*U S N F*) — Sodium Borate, 15, Sodium Bicarbonate, 15, Carbolic Acid, 3, Glycerin, 35, Water, q s to yield 1000

This has been incorporated in the *B P C*

LOTIO BORACIS — Borax, 1, Rose Water, 24

Borax, 1, Glycerin, 1, Rose Water, 16

SEILER'S ANTISEPTIC — Sodium Bicarbonate, 8 drm, Borax, 8 drm, Sodium Benzoate, 20 grains, Sodium Salicylate, 20 grains, Eucalyptol, 10 grains, Thymol, 10 grains, Menthol, 5 grains, Oil of Wintergreen, 6 minims, Glycerin, $8\frac{1}{2}$ oz, Alcohol, 2 oz, Water, to make 256 oz — *Pharm Form*

Nebula Antiseptica Alkalina — Sodium Bicarbonate, 1, Borax, 1, Sodium Benzoate, 0.04, Sodium Salicylate, 0.04, Eucalyptol, 0.02, Thymol, 0.02, Menthol, 0.01, Oil of Gaultheria, 0.01, Distilled Water, q s, to produce 100. — *B P C*

TINCTURA MYRRHÆ ET BORACIS—Myrrh, 1, Eau de Cologne, 16, Borax, 1, Water, 8, Syrup, 8

Borax, 2, Glycerin, 2, Rose Water (undiluted), 24, dissolve and add Eau de Cologne, 48 Tincture of Myrrh (*B P* 1885), 96

Tincture of Myrrh, 37 50, Oil of Bergamot, 0 20, Oil of Lemon, 0 20, Oil of Orange, 0 20, Oil of Neroli, 0 10, Oil of Rosemary, 0 20, Borax, in powder, 2 50, Glycerin, 5, Alcohol, *q s* to produce 100—*B P C*

The *B P C Supplement* alters the quantity of Tincture of Myrrh to 35, and adds 3 5 of Tincture of Krameria

TROCHISCI BORACIS—Each Lozenge contains 3 grains of Borax Sedative—*Throat*

This has been incorporated in the *B P C*

UNGUENTUM BORACIS—Borax, 1, Spermaceti Ointment, 8

For chilblains or cracked nipples

This has been incorporated in the *B P C* as follows—

Borax, in fine powder, 12 50, Spermaceti Ointment, 87 50—*B P C*

Not Official

BROMUM.

BROMINE

FR, BROME, GER., BROM, ITAL, BROMO, SPAN, BROMO.

Br, eq 79 35

A heavy, dark red liquid, which evolves dense red, intensely irritating vapours It is obtained from sea water and from some saline springs

It should be preserved in dark, amber tinted glass bottles, provided with closely fitting glass stoppers and should be kept in a cool place

Solubility—In Water, 1 in 30 by weight Readily soluble in Glycerin, Alcohol (90 p c), Ether, Chloroform, and Carbon Bisulphide, with gradual decomposition of the solvents

Medicinal Properties—Deodoriser and disinfectant Used medicinally as a sedative in the form of Bromides and Diluted Hydrobromic Acid

Official Preparations—Used to prepare Potassium Bromidum and Sodium Bromidum

Foreign Pharmacopœias—Official in Belg, Fr, Ger., Mex (Bromo), Ital, Jap, Port, Span, Swiss and U S Not in Austr, Dan, Dutch, Hung, Norw, Russ or Swed

Tests.—Bromine has a *density* of 2 97 to 3 14, and a boiling point of about 63° C (145 4° F) It is completely volatilised by heat, without leaving any residue When treated with Potassium Hydroxide Solution in excess it should form a perfectly clear liquid It this liquid be rendered faintly acid with diluted Nitric Acid it yields with Silver Nitrate Solution a yellowish precipitate soluble with difficulty in Liquor Ammonia, insoluble in Nitric Acid It gives a yellow coloration to Starch Solution, and decolorises Litmus and Indigo Solutions

The more generally occurring impurities are mineral matter, organic Bromine compounds, and Iodine Mineral matter is readily detected by a residue remaining after volatilisation Organic Bromine compounds are manifested by the failure to produce a clear solution on treatment with an excess of Potassium Hydroxide Solution Iodine by the blue colour imparted to Starch Solution by an aqueous solution of Bromine, which has been nearly decolorised by the addition of a slight excess of reduced Iron, and to which a small amount of Ferric Chloride has been added

Bromine is official in the U S P and in the F G, the former stipulates that it shall contain not less than 97 p c of pure Bromine, but gives no process for its quantitative determination.

HYPOBROMITE SOLUTION FOR UREA-ESTIMATION—Prepare a stock Solution of Soda (sp gr 1.310) by dissolving $\frac{3}{4}$ oz of pure Sodium Hydroxide in 9 fl oz of Water. To 7 fl drms of this add 42 minims (about 114 grains) of Bromine when the Solution is wanted for use.

Note—The vapour of Bromine is very irritating to the air passages. Bromine can be weighed by taking the difference between the weight of the bottle before and after pouring some out, and calculating the quantity of Soda Solution required.

Glass tubes (hermetically sealed) containing the above quantity of Bromine are made.

In place of 42 minims of Bromine, 2 fl drms of the following concentrated solution of Bromine can be used to 6 fl drms of the Soda Solution—

Liquor Bromi Conc—Bromine, 168 minims = 450 grains, Potassium Bromide, 240 grains, Water, to 1 fl oz. Mix the Bromine and Potassium Bromide and add the Water gradually with constant stirring until 1 fl oz of solution is obtained.

This has been incorporated in the *B P C* as follows—

Liquor Bromi Fortis—Bromine, by volume, $\frac{3}{4}$, Potassium Bromide, $\frac{54}{100}$; Distilled Water, *q s* to produce 100—*B P C*.

BROMIPIN—A pale yellow oily liquid. It is a Bromine addition-compound of the fatty acid of Sesame Oil, containing about 83 p c of Bromine.

Introduced for the treatment of epilepsy, in doses of 1 fl drms.

Being difficult to dispense and costly has not met with much success in epilepsy.—*L '05*, 1.710.

Prescribing Notes—It may be given in capsules, or in emulsion with Mucilage of Gum Acacia.

BROMOFORM (CHBr_3 , eq 250.96)—A heavy, translucent, colourless, mobile liquid, about twice as heavy as Chloroform. It has a pleasant ethereal odour, and a sweetish taste somewhat resembling Chloroform. The *U S P* defines Bromoform as a liquid consisting of 99 p c by weight of absolute Bromoform and 1 p c of Absolute Alcohol.

It undergoes change in colour on exposure to the light, and should therefore be kept in well stoppered dark amber-tinted glass bottles, and should be kept in a cool atmosphere.

Solubility—1 in 800 of Water, soluble in all proportions of Alcohol (90 p c), of Ether, and of Almond Oil, about 1 in 80 of Glycerin.

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c c.

Ph Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes.

Prescribing Notes—It is but slightly soluble in Water, and owing to its high sp gr, it is difficult of suspension, and from this cause accidents have occurred from patients taking an excessive quantity in the last dose of a mixture. For oral administration it is best dissolved in Almond Oil, which can then be put into capsules or made into an emulsion, see below.

It decomposes and becomes yellow on exposure to sunlight, and should not then be dispensed.

Given for the relief of whooping cough in doses of 2 to 5 drops three or four times a day, in some cases it caused languor and drowsiness, and an over-dose produced toxic symptoms.—*L '90*, 1.139, '93, 1.1062, *Pr* xlv 47, *T.G '90*, 694, '91, 214, *B M J '01*, 1.1202, 1543.

Importance emphasised of shaking the mixture containing it, before pouring out dose, and of accurately measuring it.—*B M J '07*, 1.299.

Foreign Pharmacopœias—Official in Belg, Dutch, Fr, Ger, Span, Swiss and U S.

Tests—Bromoform has a specific gravity of 2.829 to 2.883, a boiling point of 148°C to 150°C (298.4°F to 302°F), and a solidifying point of 8°C (42.8°F). A few drops of Bromoform boiled with Potassium Hydroxide Solution, and the mixture evaporated to dryness on a water-bath, yield a

residue a portion of which dissolved in Water and faintly acidified with diluted Nitric Acid yields, with Silver Nitrate, a white precipitate, insoluble in Nitric Acid, practically insoluble in Water, and a portion of the residue when dissolved in Water, and acidified with Hydrochloric Acid and Chlorine Water added yields a reddish-brown coloration, and when shaken with Chloroform the reddish-brown colour passes into the chloroformic liquid.

The more generally occurring impurities are mineral matter, free Hydrobromic Acid, Bromides, and Bromine derivatives, free Bromine, and Acetone. Mineral impurity is at once manifested by a residue remaining after evaporation. Free Acid may be detected by the reaction towards Litmus paper, of Water which has been shaken with an equal volume of the sample and allowed to separate. Bromides and Bromine derivatives by the production of a turbidity or precipitate on the addition of Silver Nitrate Solution to the same aqueous menstruum. Free Bromine is detected by the liberation of Iodine from solution of either Cadmium, Potassium, or Zinc Iodide Solution and the subsequent addition of a blue colour with Starch Solution. The U.S.P. employs the following: Starch Solution, the P.G. Zinc Iodide Solution, and the Belg. Cadmium Iodide Starch Solution. Acetone may be detected by the formation of Iodoform, when the aqueous layer separated after shaking together equal volumes of the sample and Water, is treated first with Ammonia Solution in excess and then with Iodine and Ammonium Iodide Solution. A test for this impurity is included in the U.S.P., but not in the P.G. The P.G. states that when equal parts of Bromoform and Water are shaken together, in a vessel which has been rinsed with Water, the coloration of the Acid shall take place within 10 minutes. No such test appears in the U.S.P.

Emulsio Bromoform—Bromoform 40 minims, Almond Oil, 70 minims, Gum Acacia, 40 grains, Syrup, 100 minims, Distilled Water, to 1 fl oz. Dissolve the Bromoform in the Oil or Vaseline and calculate in the usual way.

One minim is contained in 6 minims of the Emulsion.

Dose.—5 to 20 minims = 0.6 to 1.2 cc.

Mistura Bromoformi—Bromoform 12 minims, Almond Oil, 60 minims, Powdered Gum Acacia, 120 grains, Simple Syrup, 240 minims, Water, to make 3 fl oz. Dissolve the Bromoform in the Almond Oil, rub this with the Powdered Gum Acacia, add $\frac{3}{4}$ fl dr of Water and rub into a paste, gradually add the remainder of the Water, and finally the Syrup.

One minim is contained in 2 fl dr of the Mixture.

Dose.—1 to 4 fl dr = 3.6 to 14.2 cc.

BROMETHYLFORMINE (Bromaline)—A white crystalline, almost odourless powder. Has been recommended as a sedative in epilepsy.

Solubility—2 in 1 of Water, 1 in 23 of Alcohol (90 p.c.), insoluble in Ether and in Chloroform.

Dose.—5 to 30 grains = 0.32 to 2 grammes.

Tests—Bromethylformine dissolves readily in Water, forming a clear neutral solution. The aqueous solution when gently warmed with Sodium Hydroxide Solution yields on the addition of a slight excess of Iodine a strong characteristic odour of Iodoform. When Bromine Water is added in excess an orange-red precipitate is thrown down, the precipitation being more pronounced in strong solutions. When boiled with Sodium Hydroxide Solution it evolves Ammonia gas readily recognised by its odour and by its action on red Litmus paper. If the liquid be now cooled, acidified with Sulphuric Acid, and again boiled, it evolves the characteristic irritating odour of Formaldehyde. Boiled with Sodium Hydroxide Solution, cooled, and acidified with dilute Nitric Acid, it yields with Silver Nitrate Solution, a yellowish precipitate practically insoluble in Ammonia Solution, insoluble in Nitric Acid, when dissolved in Sulphuric Acid a yellowish-brown coloration, which passes into solution when shaken with Chloroform. It leaves no residue when ignited with free access of air.

Not Official BRYONIA

The Root of *Bryonia alba*, L., and of *Bryonia dioica*, Jacq

Medicinal Properties—In large doses it is an active hydragogue cathartic, in small doses it is given in pleurisy. It has also been used as a hæmostatic in menorrhagia.—L '88, if 438

It has been used for many years by the homœopaths in the form of tincture. The active principle is a glucoside

Foreign Pharmacopœias—Official in Mex and Port. Not in the others

Descriptive Notes—The root of *Bryonia dioica*, Jacq., is usually the kind met with in commerce. It occurs in circular transverse slices of a yellowish-white colour, about $1\frac{1}{2}$ to 3 inches (37 to 75 mm) or more in diameter and $\frac{1}{2}$ to $\frac{3}{4}$ inch (6 to 8 mm) thick. The narrow bark, which is a very pale brown externally, is separated from the fleshy centre by a fine line, and the cut surface is marked with concentric rings, and with radiating lines of vascular tissue. The dried root has no definite odour, but has a bitter and acrid taste. The fresh root, which is often as much as 2 feet (60 cm) long and 3 inches (75 mm) in diameter at the upper end, is occasionally offered by gardeners under the name of Mandrake root. In homœopathic medicine, the root of *Bryonia alba* is preferred, and is imported from Germany. The plant is distinguished from *Bryonia dioica* by having monocotyledonous flowers and black berries. The plant contains Brein, a glucoside not found in *B. dioica*, and the root is considered to have a different medicinal action, it is stated by Petresco to be not purgative.

TINCTURA BRYONIAE—Made from fresh Bryony Root of such a strength that 10 fl oz shall represent 1 oz of the dried root and shall contain 60 p c by volume of Alcohol.—*B P C Formulary* 1901

Fresh Bryony Root yields on an average 32 to 40 p c of dried root

Dose—1 to 10 minims = 0.06 to 0.6 c c

This has been incorporated in the *B P C*

Foreign Pharmacopœias—Mex, 1 and 5, dried Root

Antidotes—An emetic, stimulants, Brandy or Spirit of Sal Volatile

BUCHU FOLIA.

BUCHU LEAVES

NO Syn.—BUCCO, DIOSMA

FR, FEUILLES DE BUCCO, GER, BUCCOBLÄTTER

The dried leaves of *Barosma betulina*, contain a volatile oil, a bitter principle, and a mucilage

Medicinal Properties.—Tonic, stomachic, diuretic, and diaphoretic. Given chiefly in complaints of the urinary organs, as an antiseptic in chronic cystitis, and in irritation of the bladder and urethra. Also in dyspepsia, chronic rheumatism, and dropsy

Dose—Usually given in the form of Infusion or Tincture, *q v*

Official Preparations—Infusum Buchu and Tinctura Buchu

Not Official—Fluëxtractum Buchu, Infusum Buchu Concentratum, Mistura Buchu Composita

Foreign Pharmacopœias—Official in Jap, Mex, Port. and U S. Not in Austr., Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Norw, Russ, Span, Swed or Swiss

Descriptive Notes.—The dried leaves of *Barosma betulina*, Bart and Wendl, are alone official. These are easily distinguished from the leaves of *B serratifolia*, Willd and *B crenulata*, Hook, which were formerly official, and are still to be met with in commerce, by their obovate shape and recurved obtuse apex. The oil glands on the toothed margin of the leaf and its size (which is given in *B P* as $\frac{1}{2}$ to $\frac{3}{4}$ inch, or 12 to 20 mm, in length), and the layer of mucilaginous cells below the epidermis and the yellow crystals of hesperidin contained in the epidermal cells, are useful characters for distinguishing Buchu from other leaves. The oil of the leaves *B betulina* deposits crystals of Diosphenol, to which the antiseptic properties of the leaves are probably due, this is not the case with the Oil from the other species. Diosphenol has an odour like Menthol. The official leaves have also the advantage that they cannot be mistaken, like those of *B serratifolia* and *B crenulata*, for the leaves of any other species, on account of their very characteristic shape.

Tests—Buchu Leaves yield about 5 p c of ash on ignition, and 6 p c is seldom exceeded. 12 samples examined in the author's laboratory yielded from 3.6 to 5.68 p c, with an average of 4.52 p c.

Preparations

INFUSUM BUCHU. INFUSUM OF BUCHU

Buchu, 1, boiling Distilled Water, 20, infuse 15 minutes (1 in 20)

Dose.—1 to 2 fl oz = 28.4 to 56.8 c c

Not in the other Pharmacopœias

TINCTURA BUCHU TINCTURE OF BUCHU

1 of Buchu Leaves, in No 20 powder percolated with Alcohol (60 p c) to yield 5 (1 in 5)

Dose.— $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Mex, 1 and 5, both by weight. Not in the others

Tests.—Tincture of Buchu has a sp gr of 0.925 to 0.935, it contains about 4 p c w/v of total solids and about 58 p c of Absolute Alcohol

Not Official

FLUIDEXTRACTUM BUCHU—Macerate 100 of Buchu in No 60 powder with 40 of a mixture of Alcohol (95 p c) 3 and Water 1, and percolate with more of the mixture until the drug is exhausted. Reserve the first 85 of the percolate and evaporate the remainder at a temperature not exceeding 50° C. (122° F) to a soft extract, dissolve this in the reserved portion and add enough menstruum to make 100—*U S P*

INFUSUM BUCHU CONCENTRATUM—Buchu Leaves, bruised, 40, Tincture of Buchu, 22.5, Alcohol (90 p c), 10, Dilute Chloroform Water (1 in 1000), q s to make 100. Prepare by macero-expression. **Dose**—1 to 2 fl drim—*Pharm* and *Wright*, *P J* '06, 165, '07, 1621, *C D* '06, 1252, *Y B P* '07, 249. The authors state that it keeps well, but the product of dilution is not equal to the fresh infusion.

It appears in the *B P C*

MISTURA BUCHU COMPOSITA — Potassium Citrate, 30 grains, Tincture of Hyoscyamus, 30 minims, Infusion of Buchu, to 1 fl oz — *Charing Cross*

Potassium Bicarbonate, 15 grains, Tincture of Hyoscyamus, 20 minims, Spirit of Chloroform, 10 minims, Infusion of Buchu, to 1 fl oz — *King's*

Potassium Citrate, 20 grains, Tincture of Hyoscyamus, 15 minims, Infusion of Buchu, to 1 fl oz — *St Thomas's*

The last has been incorporated in the *B P C*

BUTYL-CHLORAL HYDRAS.

BUTYL CHLORAL HYDRATE

$C_4H_9Cl_2O \cdot OH_2O$, eq 191 97

White crystalline scales of a silky lustre, with a somewhat fruit-like but disagreeable odour and bitter nauseous taste

Butyl-Chloral Hydrate, formerly known as Croton-Chloral Hydrate, is, chemically, Trichlorobutylidene Glycol, and is prepared by acting upon Aldehyde or, preferably, Paraldehyde with Chlorine gas

It should be preserved in well-stoppered amber-tinted glass bottles

Solubility — 1 in 44 of Water, 1 in 1 of Glycerin (very slowly), 5 in 3 of Alcohol (90 p c), 1 in 20 of Olive Oil, 1 in 2 of Ether, 1 in 20 of Chloroform

Some books give the solubility as 4 in 1 of Glycerin, but this is incorrect

Medicinal Properties — Analgesic, is frequently but not always an efficient remedy in neuralgia of the face and head, and in tic-douloureux. As a hypnotic it is seldom used, being weaker and less certain than Chloral Hydrate

Dose. — 5 to 20 grains = 0.32 to 1.3 gramme

Prescribing Notes — *Generally given in the form of pills made with a little Compound Powder of Tragacanth and Syrup. The addition of Alcohol or Glycerin to aqueous mixtures increases its solubility*

Not Official — Mistura Butyl Chloral, Pilula Butyl-Chloral, Pilula Butyl Chloral cum Gelsemio, Syrupus Butyl-Chloral

Antidotes — The same as for Chloral Hydrate

Tests — The distinguishing tests for Butyl-Chloral Hydrate are its melting point, about $78^{\circ} C$ ($172.4^{\circ} F$), and which is officially required to be about $77.8^{\circ} C$ ($172^{\circ} F$), and its solidifying point, which should be, according to the *B P*, about $71.1^{\circ} C$ ($160^{\circ} F$). Some samples are acid, very pungent and acrid. Of these the author found that 1 gramme heated in a porcelain capsule over a water-bath for 10 minutes wholly volatilised, but the sample lost its pungency and acidity after having been washed with about twice its weight of Water, pressed, and dried by exposure to air, and when heated as above lost less than half its weight. The slow volatility of a sample may therefore be taken as a test of purity. An acrid sample by washing and drying had its melting point raised from $73.9^{\circ} C$ ($165^{\circ} F$) to $78.9^{\circ} C$ ($174^{\circ} F$). When warmed with concentrated Sulphuric

Acid, Trichlorobutyl-aldehyde separates out in oily drops. The aqueous solution of Butyl-Chloral Hydrate reduces Silver Ammonio-nitrate Solution.

The more generally occurring impurities are free acid, Chloral Hydrate, and Chlorine derivatives. The behaviour of the aqueous solution towards blue Litmus paper affords a measure of the free acid; the non-production of a turbidity with Silver Nitrate Solution indicates the absence of free Hydrochloric Acid or Chlorine derivatives, and should the sample evolve no odour of Chloroform when heated with Calcium Potassium, or Sodium Hydroxide Solution, the absence of Chloral Hydrate may be inferred.

Its behaviour when warmed with concentrated Sulphuric Acid affords a test for Chlorine derivatives. Thus heated the liquid should not turn brown. The *BP* says nothing respecting these Chlorine derivatives, only testing for freedom from acidity and Chloral Hydrate, the aqueous solution is required to be neutral or but slightly acid to Litmus, the salt should leave no weighable residue when ignited with free access of air.

Not Official.

MISTURA BUTYL-CHLORAL — Butyl-Chloral Hydrate, 5 grains, Glycerin, 15 minims, Chloroform Water, $\frac{1}{2}$ fl oz, Water, to 1 fl oz.

This has been incorporated in *BP C* as follows —

Butyl Chloral Hydrate, $4\frac{1}{2}$ grains, Glycerin 15 minims, Chloroform Water, 240 minims, Distilled Water, *q s* to make 1 fl oz — *BP C*

PILULA BUTYL-CHLORAL — Butyl-Chloral Hydrate, 5 grains, Compound Powder of Tragacanth, 1 grain, Syrup *q s*, in 1 pill.

PILULA BUTYL-CHLORAL CUM GELSEMIO — Butyl-Chloral Hydrate, 3 grains, Alcoholic Extract of Gelsemium, 1 grain — *Guy's and Sheffield Union*

PILULA BUTYL-CHLORAL ET GELSEMINÆ — Butyl-Chloral Hydrate, 3 grains, Gelseminæ Hydrochloridum, $\frac{1}{100}$ grain, Pulvis Tragacanthæ Compositus, 1 grain, Syrupi Glucosi *q s* — *Westminster*

SYRUPUS BUTYL-CHLORAL — Butyl-Chloral Hydrate, 16 grains, Syrup, *q s* to make 1 fl oz. Dissolve by aid of heat — *BP C*

This has been incorporated in *BP C*

Dose — 1 to 4 fl drms = 3.6 to 14.2 cc

Not Official

BYNE. MALT.

Good Malted Barley is tolerably uniform in diastase, and the widely differing results published from time to time by different analysts as to the strength of commercial Extracts must be due partly to a deviation or diastase in the manufacture of the Extracts, and partly to an ambiguity attaching to the phrase 'conversion of Starch'.

EXTRACTUM BYNES *Syn* EXTRACTUM MALTUM MALT EXTRACT.

Is made by infusing or mashing ground Malt in Water at a temperature under 160° F, preferably 140° F, filtering and evaporating the solution *in vacuo* to the consistence of a thick syrup. It is more convenient to use when it is evaporated only to a thin syrup, but in that condition the Extract is more liable to undergo fermentation.

Medicinal Properties—Malt Extract is prescribed as a nutrient in wasting diseases, and where the digestion is weak it is given for its diastase value of converting Starch into Maltose and Dextrin. It is also given with Cod Liver Oil. It is useful for covering the taste of nauseous drugs.

Dose—A teaspoonful to a tablespoonful, immediately after food.

Hale White states, 'like the ferments of pancreatic juice and saliva, diastase can only act in an alkaline medium, and therefore extract of malt should not be given till at least two hours after a meal'—*Hale White Mat Med*. But the usual custom of most prescribers is probably that expressed in the following reasoned statement of the *B M J* '08, 1 363—

Malt Extract is prescribed with one of two objects (1) To increase the supply of carbohydrates in the diet. When given for this purpose it is a matter of indifference at what hour it is taken, and the choice may well be left to the individual taste of the patient. (2) To help in the conversion of starch into sugar by means of the diastase it contains. The contents of the pyloric half of the stomach become acid at a very early stage in digestion. Those of the cardiac half are alkaline for a considerable period, so that amylolytic digestion may continue in the latter after a full meal for an hour or more. The first part of a meal finds its way into the pyloric half, the second part remains in the cardiac half until the former passes into the duodenum. Hence if the latter part of a meal is largely starchy, malt extract taken with it or immediately after it helps in the conversion of starch into sugar. If taken an hour or more after the meal the Malt will be useless, as the contents of all parts of the stomach are then acid.

It is very useful when mixed with baked wheaten flour to form foods for infants and invalids when a certain amount of pre digestion is required.

For a substance with similar properties see TAKA DIASTASE.

Foreign Pharmacopœias—The *U S P* 1882 ordered the Malt to be macerated in cold Water for six hours, then digested for an hour at 181° F, strained and evaporated at a temperature not exceeding 181° F to the consistence of Honey. This contained active diastase. It was omitted in *U S* 1893. It was re introduced in *U S P* 1905 as follows—

Extractum Malti.—Upon 1000 grammes of Malt in coarse powder (not finer than No 12), contained in a suitable vessel, pour 1000 c c of Water, and macerate for six hours. Then add 4000 c c of Water, heated to about 80° C (86° F), and digest for an hour at a temperature not exceeding 55° C (131° F). Strain the mixture with strong expression. Finally, by means of a water-bath, or vacuum apparatus, at a temperature not exceeding 55° C (131° F), evaporate the strained liquid rapidly to the consistence of thick Honey.

This has been incorporated in the *B P C*.

***Test**—For Malt or Malt Extract, three solutions, *A*, *B*, and *C*, are required. (*A*) Infuse 5 grammes of ground Malt in 100 c c of Water at 140° F (60° C) for one hour, cool to 60° F (15 5° C), and make up to 100 c c with Water, filter. For testing Malt Extract, dissolve 5 grammes of the Extract in sufficient Water to make 100 c c of solution. (*B*) Mix 1 gramme of Potato Starch with 10 c c of Water, add to it 90 c c of boiling Water, boil the mixture for ten minutes, cool to 60° F (15 5° C) and make up to 100 c c, strain through fine muslin. (*C*) Dilute 1 c c of *B P* Volumetric Solution of Iodine to 75 c c with Water.

Method—Run 2 c c of the Iodine Solution into each of one dozen test-tubes. Bring solution *A* and solution *B* to 100° F (37 8° C), place 50 c c of *B* in a beaker immersed in Water at 100° F (37 8° C), and add to it 10 c c of *A*, at the end of a minute draw off 2 c c of the mixture and add it to the Iodine Solution in one of the test-tubes, and at the end of each subsequent minute repeat the operation. If the test-tubes are arranged in the order in which the solution is added, the colour in each test-tube will represent the amount of action in a given time represented by minutes. As it occupies from ten to fifteen seconds to run the Malt Solution from a pipette into the Starch, we usually start the stop-watch or chronograph when half of the solution has run out of the pipette. When a first-class sample of Malt Extract is used, the contents of the first test-tube will be of a blue colour, the second will be red, and the third or fourth yellow, but the changes will be somewhat slower in a sample which is not so good.

Six of the best known brands of Malt Extract examined by this test ceased to produce a red colour at the end of three, five, ten, fifteen, and fifteen minutes respectively, showing a variation of about ten to fifteen minutes, in the digestion of *their own weight* of Starch. A fluid Malt Extract, containing Alcohol, ceased to give a red colour at the end of thirty-five minutes.

The best sample, when treated with *five times* its weight of Starch, ceased to produce a red colour at the end of fourteen minutes.

It is important that the conditions should be the same in each experiment, for any variation in the quantity of Starch Solution or the volume of liquid employed will affect the results, but under the conditions given, when the colours are viewed in series, two independent workers should not vary more than 1 minute in the reading.

This process has been found (*Suggested Standards of Purity for Food and Drugs*, p. 184) useful and convenient, but it is mentioned that it may have been better to adopt a standard time and vary the proportion of Starch. This suggestion would render the process much less convenient, and the amount of Starch digested in a Standard time can easily be calculated by a simple formula, the activity of Malt Diastase towards Starch Solution, unlike Pancreatic Diastase (Ankylopilin), being inversely proportional to the amount of enzyme present. A more striking contrast is, moreover, obtained by taking the relative time required by different samples to digest a definite weight of Starch.

LIQUID MALT—Malt Extract, sp gr 1.375, to which diluted Alcohol is added, sufficient to produce a liquid, sp gr 1.250, containing 73 p.c. of Alcohol (90 p.c.) by weight, equal to about 15 p.c. Proof Spirit.

Preparations somewhat similar to this are sold as **Fluid Extract of Malt**, **Bynn**, etc.

Liquid Malt Extract—Extract of Malt (sp gr. 1.375), by volume, 68, Alcohol (90 p.c.), 7.50. Distilled Water, q.s. to produce 100.—Mix to form a liquid sp gr 1.2—**B.P.C.**

FLUIDEXTRACTUM MALTI (U.S.N.F.)—Malt in coarse powder, 100, percolated with a mixture of Alcohol (94 p.c.) 1 and Water 3, until it percolate weighs 75.

MALT EXTRACT WITH COD LIVER OIL—This is supplied under several well-known brands, but can be prepared extremely easily by turning ordinary Malt Extract with 10 to 15 p.c. of V.C. oil, and heating to 120° F., adding the oil and shaking thoroughly until mixed. The commercial product contains from 20 to 30 p.c. of Cod Liver Oil.

Examination of commercial samples gave from 20 to 30 p.c. of Oil by volume.—*P.J.* (3) xxv 162.

Prescribing Note—*Usually given in milk*

EXTRACTUM MALTI CUM OLEO MORRHUÆ (B.P.C.)—Extract of Malt, 17 fl oz, Cod Liver Oil, 3 fl oz. Heat the Extract to 110° F., and pour it into a warm mortar, add the Oil gradually and with constant trituration.—*B.P.C. Formulary* 1901.

B.P.C. incorporates this formula without heating the extract.

EXTRACTUM MALTI FERRATUM—Iron Pyrophosphate, 2, Water 3. Dissolve and add Extract of Malt, q.s.

Dose.—1 to 4 drms = 3.6 to 14.2 cc.

Each fl drms contains about 1 grain Iron Pyrophosphate.

TAKA-DIASE—A powder of a light brown colour, possessing a nutty taste. Derived from a fungus of the species *Eurotium Oryzae*. It possesses high diastasic properties, readily converting over a hundred times its weight of Starch at body temperature.

Specially indicated as an artificial digestant of starchy food in dyspepsia cases in which there is a deficiency of saliva. Also found useful in hyperacidity of the stomach and in gouty dyspepsia.—*L* '96, 1856, '03, n. 1052, *A.J.P.* 188.

Most useful in gouty dyspepsia, it encourages the digestion of carbohydrates, prevents the development of fatty acids which, by their irritating effects,

are so common a factor in the development of gouty dyspepsia, a $2\frac{1}{2}$ grain tablet before each meal —Pr '07, 1 168

Dose—5 to 10 grains = 0.32 to 0.65 gramme, in capsules in the middle of or immediately after a meal

Diastase from Malt is official in Fr, Jap, Mex and Span

CADINUM OLEUM.

OIL OF CADE

B P Syn. JUNIPER TAR OIL

Fr, HUILE DE CADE, Ger, KADEOL, Ital, OLIO CADINO, Span, BREA DE OXICEDRO

A brownish or dark brown, viscid fluid, of an oily nature, with a tarry odour and an empyreumatic and somewhat bitter taste; a product of the dry distillation of the Branches and Wood of *Juniperus Oxycedrus*, L, and other species. It contains a high percentage of the sesquiterpene, Cadinene

Solubility—Mixes in all proportions with Chloroform and Ether, partially soluble in Alcohol (90 p c), slightly soluble in Water

Medicinal Properties—An agreeable form of Tar. Used as a stimulant in cases of psoriasis and of chronic eczema

Prescribing Notes—It is used in the form of Ointment, the Oil mixed with equal parts of Yellow Wax, and further diluted with Lard or Vaseline if required

Foreign Pharmacopœias—Official in Austr, Dan, Dutch, Hung., Ital, Norw, Port, Span (Brea de Oxicedro), Russ, Swed (Pyrooleum Oxycedri), Swiss and U S. Not in the others

In **Balzer's Cade Baths** the oil is emulsified before being added to the bath

An emulsion of Oil of Cade and a fluid extract from the decoction of the tops of the Tannus Pines. The baths are given daily, and can be used in all varieties of psoriasis

Cade Bath Fluid is put up in bottles, one of which, being mixed with a little warm Water, is added to an ordinary bath, which should last from half an hour to an hour, and should be accompanied by slight friction on the affected patches

These baths may be used as an adjuvant to the ordinary treatment, and are suitable both for hospital and private cases

Tests—Oil of Cade has a specific gravity of about 0.990. When shaken with Water and filtered it should yield an almost colourless filtrate possessing an acid reaction. In a sample (sp gr 0.996), examined by the author, the acidity amounted to 0.7 p c of pure Acetic Acid

The filtered aqueous solution should yield a red coloration with diluted Ferric Chloride Test-solution

The B P states that the filtered aqueous solution is almost colourless and possesses an acid reaction, and the U S P that it imparts to Water an acid reaction

VASOLIMENTUM EMPYREUMATICUM, — Juniper Tar Oil, 25, Vasoliment, 75 —Hager.

Parogenum Empyreumaticum.—Oil of Cade, 25; Parogen, 75 —B P C

CAFFEINA.

CAFFEINE

B P Syn — THEINE

N O Syn — COFFEINA

FR, CAFFEINE, GER, KOFFEIN, ITAL, CAFFEINA, SPAN, CAFEINA



Fine white silky acicular crystals, odourless, and possessing a bitter taste

A feebly basic alkaloid, contained in the prepared and dried Leaves of *Camellia Thea*, Link, the dried seed of *Coffea Arabica*, L, and also in other plants. It is chemically allied to Theobromine, being Methyl-theobromine (Trimethyl-xanthine)

The quantities yielded are about as follows. Tea Leaves, 3 to 4 p c, Coffee Seeds, 1 p c, Guarana, 5 p c, Maté or Paraguay Tea, 1 p c, Kola Nut, 2 to 3 p c.

Solubility—1 in 68 of Water, 1 in 40 of Alcohol (90 p c); 1 in 7 of Chloroform, 1 in 400 of Ether, 1 in 1 of boiling Water.

Medicinal Properties.—A valuable heart tonic and diuretic, especially in cases of loss of compensation with cardiac dropsy. To be given with caution in the presence of renal disease.

Given in 1 grain doses every hour for migraine and hemiplegia, also in the form of Effervescent Caffeine Citrate (1 grain in each dram)

It is eliminated but slowly by the kidneys, and its action on the heart is cumulative.—B M J E '00, 1 35

Used by malingering soldiers to produce symptoms of cardiac disease—L '00, 1 1406

Diuretic action of Caffeine—'01, 11 7

Specially valuable in nervous affections. In 5-grain doses every four hours.—Pr '11, 318

Dose.—1 to 5 grains = 0.06 to 0.32 gramme

Ph. Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes

Prescribing Notes.—Given in cachets, in mixtures, or in pills made with 'Diluted Glucose', also in the form of effervescent preparations. For hypodermic use 20 grains can be dissolved in 60 minims of Water on the addition of 25 grains of Sodium Salicylate, or 25 grains of Sodium Benzoate. See also Caffeine Sodio-Salicylas, and Caffeine Sodio-Benzoylas.

Official Preparations.—Caffeine Citras, Caffeine Citras Effervescent

Not Official.—Caffeine Hydrobromidum, Caffeine Diiodo-Hydrobromidum, Caffeine Sodio-Benzoylas, Caffeine Sodio-Salicylas, Caffeine Valerianas, Ethoxycaffeinum

Foreign Pharmacopœias.—Official in Austr, Belg, Dan, Dutch, Fr, Ger (Koffein), Hung, Ital (Caffeina), Jap, Mex, Port, Russ, Span (Cafeina), Swed, Swiss and U S. Not in Norw

Tests.—The distinguishing test for Caffeine is the Murexide test which consists in dissolving the alkaloid in a small quantity (about 1 to 1 c c) of concentrated Hydrochloric Acid, adding a crystal of Potassium Chlorate and evaporating the mixture to dryness on a water-bath. When the residue is subjected to the action of Ammonia

gas or is moistened with a little Ammonia Solution, a fine purple coloration is produced. The colour is not affected by an excess of Ammonia, but is immediately discharged by a fixed alkali. A similar coloration is produced when the test is performed with Chlorine or Bromine Water in place of Potassium Chlorate and Hydrochloric Acid, Chlorine Water is used in the *PG* test. The usual alkaloidal reagents precipitate Caffeine only imperfectly. Neither Mercuric-potassium Iodide (Mayer's) Solution nor Iodo-potassium Iodide (Wagner's) Solution precipitate Caffeine from neutral solutions, this non-precipitation serving to distinguish Caffeine from other official alkaloids. If the Iodo-potassium Iodide Solution is preceded or followed by the addition of some dilute mineral acid, a dark-reddish precipitate is at once thrown down. Upon this reaction is based (*CN* '97, 99) a method for the determination of Caffeine. Tannic Acid produces a white precipitate in moderately dilute solutions, but the extent of the reaction is largely dependent on the temperature.

Caffeine is completely extracted by Chloroform from a slightly acid or slightly ammoniacal aqueous solution, and is the solvent generally employed in its gravimetric determination. It dissolves without colour in Sulphuric and in Nitric Acids. It melts, when anhydrous at 231.5°C (448.7°F). No melting point is given in the *BP*. The *PG* gives 230.5°C (446.9°F), and the *USP* 236.8°C (458.3°F) after drying till constant in weight at 100°C (212°F). Theoretically, it should contain 8.49 p.c. of Water, and the *BP* states that the crystals lose this amount at 100°C (212°F), but commercial Caffeine generally loses about 7 p.c. on drying. It has been pointed out (*PJ* '00, u 148) that the Pharmacopœia errs in its method of expression, as commercial Caffeine probably never contains 8.49 p.c. of Water, owing to the facility with which the crystals effloresce. The *PG* states that in the air Caffeine loses 1 part of its Water of crystallisation and at 100°C (212°F) it becomes anhydrous. The sublimation point is given in the *PG* as 180°C (356°F), in the *USP* as 178°C (352.4°F). The *BP* says that 'at a higher temperature than 100°C (212°F) it melts and volatilises without decomposition'. It has been shown (*PJ* (3) xxiii 213) that Caffeine which has been dried at the ordinary temperature over Sulphuric Acid till constant in weight undergoes no further material loss on prolonged exposure in an open dish in the water oven at 100°C (212°F), and that it does not volatilise with steam during the evaporation of its solutions.

The more generally occurring impurities are alkaloids other than Caffeine, organic impurities and mineral matter. The presence of alkaloids other than Caffeine is shown by the formation of a precipitate with Mercuric-potassium Iodide (Mayer's) Solution or Iodo-potassium Iodide Solution, organic impurities by the production of coloured solutions in Sulphuric Acid, and mineral matter by a residue remaining after sublimation.

Potassium Chlorate and Hydrochloric Acid—A small quantity of Caffeine is dissolved in 1 c.c. of Hydrochloric Acid in a porcelain dish, a little Potassium Chlorate is added and the whole evaporated to dryness on a water-

bath, the dish is then inverted over a vessel containing a few drops of Ammonia Water, the residue acquires a milky colour which is destroyed by fixed alkalis *U S P*. A similar test is given in the *P G*, but Chlorine Water is used instead of Potassium Chlorate and Hydrochloric Acid. One part of Caffeine with 10 parts of Chlorine Water are evaporated on a water-bath and yield a yellowish-red residue, which with a little Ammonia Solution becomes coloured a rich purple. It also evaporates an indefinite quantity of Caffeine with Potassium Chlorate and a few drops of Hydrochloric Acid.

Potassium Dichromate and Sulphuric Acid—If a fragment of Caffeine be dissolved in Sulphuric Acid, and a minute fragment of Potassium Dichromate be added to the liquid, a yellowish-green colour which gradually becomes green will be produced, *U S P*.

Chlorine Water—A cold saturated aqueous solution of Caffeine should not become turbid with Chlorine Water, *P G*.

Iodine Solution—A cold saturated aqueous solution should not become turbid with Iodine Solution *P G*.

Ammonia Solution—A cold saturated aqueous solution should not become coloured on the addition of Ammonia Solution, *P G*.

Sulphuric Acid or Nitric Acid—*P G* and *U S P* require that Caffeine should dissolve without coloration in Sulphuric Acid or in Nitric Acid. The *P G* uses 0.1 gramme of Caffeine and 1 c.c. of either Acid.

Preparations

CAFFEINÆ CITRAS. CAFFEINE CITRATE

An unstable combination of Caffeine and Citric Acid, which readily undergoes dissociation in the presence of Water.

A fine white, odourless powder possessing an acid and somewhat bitter taste prepared by stirring one part of Caffeine into a heated solution of 1 part of Citric Acid and 2 parts of Distilled Water, the mixture being subsequently evaporated to dryness on a water-bath and kept constantly stirred towards the end of the evaporation. It is then reduced to powder.

Solubility—1 in 32 of Water, 1 in 22 of Alcohol (90 p.c.), 1 in 10 of a mixture of 2 parts Chloroform with 1 part Alcohol (90 p.c.).

When combined with Sodium Benzoate, it acted rapidly and efficaciously in a case of transudic incompetency—*M P* '04, ii 515.

Dose.—2 to 10 grains = 0.13 to 0.65 gramme.

Foreign Pharmacopœias—Official in Hung, Mex, Span (*Citrato Caffeico*), Swiss and U.S. Not in the others.

Tests.—Caffeine Citrate should afford the reactions distinctive of Caffeine appearing in the large type under that substance. The aqueous solution, after neutralisation and separation of the Caffeine by shaking with Chloroform, yields with Calcium Chloride Solution a white precipitate insoluble in Potassium Hydroxide Solution, with Silver Nitrate Solution a white precipitate soluble in Ammonia Solution, which is distinguished from Tartrate by yielding no mirror on boiling. The salt is official in *BP* and *U S P*, but not in *P G*. Neither *BP* nor *U S P* gives any process for ensuring the presence of the requisite proportion of Caffeine. It should contain not less than 50 p.c. by weight of Caffeine as gravimetrically determined, by rendering an aqueous solution of the salt faintly alkaline.

with Potassium or Sodium Hydroxide Solution and shaking out the Caffeine with successive quantities of Chloroform. The mixed chloroformic solutions are evaporated to dryness and the residue dried till constant in weight at 100° C (212° F) and weighed. The Swiss Pharmacopœia states that it shall contain not less than 50 p c nor more than 75 p c of Caffeine.

The Citric Acid may be determined by direct titration with Normal Volumetric Potassium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. Caffeine is so feebly basic that it exercises no influence on the indicator. Theoretically it should contain about 50 p c of Citric Acid.

The *USP* states that 1 part of Caffeine forms a clear syrupy solution with about 4 parts of hot Water, which is correct, and agrees with the Swiss Pharmacopœia statement, 'soluble' in 4 parts of hot Water. On dilution with 5 parts of Water a white crystalline precipitate separates which redissolves when about 25 parts of Water have been added. The *BP* states that 'with 3 parts of Water it forms a clear syrupy solution,' which is incorrect, but more Water dissociates the salt and affords a white precipitate of Caffeine, which redissolves when excess of Water is added. It has been pointed out (*PJ* '00, ii 148) that in reality with 3 parts of Water it forms a stiff paste and only forms a solution when the mixture is gently warmed, but, on cooling, it again forms an almost solid mass of acicular crystals. On adding a little Water to the warm solution the Caffeine separates out as a mass of acicular crystals and not in the form which might be described as a 'white precipitate'.

The more generally occurring impurities are Calcium, Lead, Chlorides, Sulphates, Tartrates, and mineral matter. The presence of Calcium may be detected by Ammonium Oxalate Solution, Lead by Hydrogen Sulphide Solution in a solution made faintly acid with Hydrochloric Acid, Chlorides by Silver Nitrate Solution, Sulphates by Barium Chloride or Nitrate Solution. The *USP* tests for Tartaric Acid by heating 0.25 gramme of the salt for fifteen minutes on a water-bath with 5 c c of concentrated Sulphuric Acid, care being taken to protect the mixture from dust, no brown or black coloration should be developed, but only a lemon-yellow colour. The *BP* allows 'a mere trace of ash when heated in air'.

The Belgian Pharmacopœia states that when Caffeine Citrate is prescribed for internal use, the indicated dose should be dispensed as a mixture of equal parts of Caffeine and of Citric Acid.

CAFFEINÆ CITRAS EFFERVESCENS EFFERVESCENT CAFFEINE CITRATE

Contains about 2 p c of Caffeine (4 p c of Caffeine Citrate), or nearly 9 grains of Caffeine in the oz.

Dose.—60 to 120 grains = 4 to 8 grammes

Foreign Pharmacopœias—Official in *US*, containing 2 p c of Caffeine, and in *Spain*. Not in the others.

For other Caffeine Citrate Effervescent Compounds, see *PHENACETIN* and *ASAACETONE*.

more than Caffeine scented with Valerianic Acid, but for all purposes of practical dispensing, a product, obtained by absorbing 1 of anhydrous Valerianic Acid by 4 of anhydrous Caffeine, is superior to anything commercially obtainable, and when assayed by the above method shows when freshly prepared about 17 p c, and even after keeping a considerable time, about 9.7 p c of anhydrous Valerianic Acid. The residue obtained on washing and evaporating the chloroformic solution (after titration) to dryness should answer the tests distinctive of Caffeine given under that substance.

ÆTHOXYCAFFEINUM—A compound of Caffeine, containing an additional Æthoxyl group. Colourless, crystalline needles, less soluble in Water than Caffeine, readily soluble in Alcohol. Heart tonic and diuretic, also narcotic. Given with Sodium Salicylate in migraine and trigeminal neuralgia. Simultaneously it acts as an anæsthetic. It readily forms soluble double salts with Sodium Benzoate and Salicylate.

Dose—1 to 3 grains = 0.06 to 0.2 gramme

The following have also received notice in Medical Literature—Caffeine-chloral, Caffeine Tri-bromide, Caffeine Sodium Iodide, Caffeine Sulphate, Caffeine Vanadate, Symphoral L (Lithium Caffeine Sulphonate), Symphoral N (Sodium Caffeine Sulphonate), and Symphoral Sr (Strontium Caffeine Sulphonate).

CAJUPUTI OLEUM.

OIL OF CAJUPUT

FR, HUILE VOLATILE DE CAJEPUT, GER, CAJEPUTOL, ITAL, ESSENZA DE CAJEPUT, SPAN, ESENCIA DE CAJEPUT

A transparent green or bluish-green, thin, oily liquid, possessing an agreeable odour resembling Eucalyptus and a pungent, camphoraceous taste.

A volatile Oil distilled from the leaves of *Melaleuca Leucadendron*.

Solubility—In all proportions of Alcohol (90 p c)

Medicinal Properties.—Antispasmodic, carminative and stomachic. Counter-irritant. Given in flatulent colic, hysteria, and other spasmodic and nervous affections. Externally, diluted with Olive Oil (1 to 2), or with Linimentum Terebinthinæ, it is used for chilblains and to allay rheumatic and gouty pains. Applied on lint for toothache.

Dose— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c c

Prescribing Notes—Given on Sugar, or in Pill or in the form of Spirit of Cajuput. Occasionally as much as 10 minims are given in a mixture. See below.

Official Preparations—Spiritus Cajuputi, contained in Linimentum Crotonis.

Not Official—Mistura Cajuputi.

Foreign Pharmacopœias—Official in Austr, Dutch, Ital, Jap, Norw, Port, Russ, Span, Swed, Swiss and U S. Not in Belg, Dan, Fl, Ger., Hung, or Mex.

Tests.—Cajuput Oil has a specific gravity of 0.919 to 0.930. The *USP* gives 0.915 to 0.925 at 25° C. (77° F.). The optical rotation is not given in the *BP*, according to the *USP* it should not exceed -2° in a 100 mm tube at a temperature of 25° C (77° F.). Oils are occasionally met with having as high a rotation as $-3^{\circ} 40'$.

It should contain about 55 to 65 p c of Cineol (Eucalyptol) as determined by the Phosphoric Acid process described under Eucalyptus Oil. The *B P* makes no attempt to determine the actual amount of Cineol, only requiring that the Oil shall become semi solid when stirred with one-third or one half its volume of Phosphoric Acid, specific gravity 1.750, the mixture being kept cool. The *USP* measures the actual amount of Cineol produced on decomposing the Cineol Phosphate. A measured quantity of 10 c c of the oil is dissolved in 5 times its volume of purified Petroleum Benzin, cooled by a freezing mixture, Phosphoric Acid, specific gravity 1.707, cautiously added until the white crystalline magma commences to assume a yellowish or pinkish tint. The crystals are filtered under pressure, washed with cold purified Petroleum Benzin and dried by pressure between porous plates. The crystals are decomposed by warm Water and the volume of Cineol read off in a narrow graduated cylinder, the oil should yield not less than 55 p c.

The green colour of the oil is generally supposed to be due to Copper, which may be detected by shaking the oil with an equal volume of Water containing a drop or two of diluted Hydrochloric Acid. The oil loses its green colour and the aqueous portion when tested with a drop or two of Potassium Ferrocyanide Solution, yields a reddish-brown colour if Copper be present. The *USP* includes a test for absence of Copper, which is given below in small type under the heading of Potassium Ferrocyanide.

Potassium Ferrocyanide—On shaking 5 c c of the Oil with 5 c c of Water containing 1 drop of diluted Hydrochloric Acid, a reddish-brown colour should not be produced in the acid liquid when separated from the Oil, if a drop of Potassium Ferrocyanide Test Solution be added (absence of Copper).

Preparation

SPIRITUS CAJUPUTI SPIRIT OF CAJUPUT

Oil of Cajuput, 1, Alcohol (90 p c), *q s* to yield 10

Dose —5 to 20 minims = 0.3 to 1.2 c c

Not Official

MISTURA CAJUPUTI—Oil of Cajuput, 80 minims, Compound Spirit of Orange (*USP*), 80 minims, Powdered Tragacanth, 8 grains, powdered Gum Acacia, 120 grains, Glycerin, 1 fl oz, Chloroform Water, to 8 oz

Dose —1 to 4 fl drms = 3.6 to 14.2 c c

Not Official

CALAMINA PRÆPARATA

PREPARED CALAMINE

Native Zinc Carbonate, calcined in a covered earthenware crucible at a moderate temperature, powdered and freed from gritty particles by elutriation. Genuine Calamine has a yellowish grey colour, the reddish varieties are generally made on a basis of Barium Sulphate.

Medicinal Properties—Mildly astringent, used in face lotions and dusting powders.

Calamina Artificialis.—Dissolve 861 parts of Crystallised Sulphate of Zinc in Water, and add 15 fluid parts of Strong Solution of Perchloride of

Iron (*B P*) Next dissolve 890 parts of Crystallised Sodium Carbonate in a separate quantity of Water, mix the solutions. Shake well, collect the precipitate on a filter, wash until free from Sulphate, well drain, heat in a light tin over a fire until a portion of it ceases to effervesce on the addition of an acid. Grind to an impalpable powder—*Y B P* 1893, 210, *P J* '98, 1 622, *C D* '98, 1 73

Calamina Facititia (*B P C*) uses the above process, with quantities as follows—Zinc Sulphate, 56, Sodium Carbonate, 58, Strong Solution of Ferric Chloride, 1, Distilled Water, *q s*

LINIMENTUM CALAMINÆ—Prepared Calamine, 20 grains, Zinc Oxide, 15 grains, Solution of Lime, 2 dim, Water, 2 dim, Olive Oil, to 1 fl oz—*Guy's*

Variations of this are given in Great Northern, Middlesex and University Hospital Pharmacopœias. The following is sometimes prescribed, but unless treated in the manner here described it is difficult to dispense—Calamine, 3 dim, Zinc Oxide, 2 dim, Lime Water, 4 fl oz, Olive Oil, 4 fl oz

Rub the powders with the Lime Water in a mortar to a smooth cream, and then add the whole of the Oil at once and stir together

This has been incorporated in the *B P C*
Prepared Calamine, 4 50, Zinc Oxide, 3 50, Solution of Lime, 50, Olive Oil, *q s* to produce 100—*B P C*

LOTIO ZINCI OXIDI—Zinc Oxide, 60 grains, Prepared Calamine, 60 grains, Glycerin, 60 minims, Water, 1 fl oz

A mild astringent in chronic eczema and acne rosacea

Most of the Hospital Formulary give a formula under the heading **Lotion Calaminæ** or **Lotio Zinc Oxidi**

LOTIO CALAMINÆ—Levigated Calamine, 40 grains, Zinc Oxide, 20 grains, Glycerin, 20 minims, Water (or Rose Water), *q s* to make 1 fl oz. Elutriate the Calamine and Zinc Oxide by titrating with the Water and decanting from the siliceous matter, then add the Glycerin—*Canadian Formulary* (first issue)

This has been incorporated by the *B P C* as follows—

Prepared Calamine, 10, Zinc Oxide, 5, Glycerin, 5, Rose Water, *q s* to produce 100

Lotio Calaminæ—Levigated Calamine, 40 grains, Zinc Oxide, 20 grains, Glycerin, 20 minims, Lime Water, *q s* to make 1 fl oz—*Canadian Formulary* 1908

UNGUENTUM CALAMINÆ—Prepared Calamine, 1, Benzoated Lard, 5—*B P* 1885

This has been incorporated in the *B P C*

CERATUM CALAMINÆ (*Ph Lond*)—Calamine, 7½, Beeswax, 7½, Olive Oil, 20

This has been incorporated in the *B P C* as follows—

Ceratum Calaminæ—Prepared Calamine, 2, Yellow Beeswax, 2, Olive Oil, 6

Ph Edin—Calamine, 1, Simple Cerate, 5, commonly known as *Turner's Cerate*

CALCII CARBONAS PRÆCIPITATUS.

PRECIPITATED CALCIUM CARBONATE

B P Sym—PRECIPITATED CHALK

FR, CARBONATO DI CALCIO PRÉCIPITÉ, GR, CALCIUMCARBONAT,
ITAL, CARBONATO DI CALCIO, SPAN, CALCIÓCARBONATO

CaCO_3 , eq 99 26

A white, tasteless powder

It may be obtained by precipitating a soluble Calcium salt, usually the Chloride with Sodium Carbonate

Solubility—Insoluble in Water and Alcohol Soluble with effervescence in dilute mineral Acids and some of the organic Acids

Medicinal Properties—Antacid, astringent and desiccant Used in dyspepsia with acidity, valuable in diarrhoea, as a dusting powder in eczema, and for burns

Dose—10 to 60 grains = 0.65 to 4 grammes

Official Preparation—Used in the preparation of Trochiscus Bismuthi Compositus

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests—Precipitated Chalk effervesces upon the addition of acids, yielding a colourless and odourless gas, which affords, when passed into Lime Water or into Barium Hydroxide Solution, a white precipitate

When dissolved in just sufficient Hydrochloric Acid to effect solution, boiled and cooled, it answers the tests distinctive of Calcium, which are that, when mixed with sufficient Ammonia Solution to render the solution faintly alkaline, Ammonium Oxalate Solution produces a white precipitate, soluble in Hydrochloric Acid, insoluble in Acetic Acid, Ammonium Carbonate Solution produces a white precipitate which, after well boiling, is insoluble in Ammonium Chloride Solution

The *U S P* states that the salt should contain not less than 99 p c of pure Calcium Carbonate, but no process is given for the determination The *B P* and *P G* neither state the requisite percentage nor method of determination It is usually determined gravimetrically as Oxalate, the Calcium Oxalate being ignited and weighed as Calcium Oxide

The more generally occurring impurities are Aluminium, Iron, Magnesium, Chlorides, Phosphates and Sulphates Aluminium, Iron and Phosphates, if present, may be detected by Ammonia Solution in presence of Ammonium Chloride The *P G* gives a specific test for Iron with Potassium Ferrocyanide Solution, which is described in small type below The *U S P* includes a test for heavy metals The standard suggested (*C D* '08, 1796) for Lead is 10 parts per 1,000,000, and for Arsenic 5 parts per 1,000,000 Both *P G* and *U S P* fix a limit of soluble impurities, ascertained by evaporating an aqueous extract Chlorides and Sulphates are indicated by Silver Nitrate and Barium Chloride or Nitrate Solutions respectively

Ammonia—If to 20 c c of the Hydrochloric Acid solution, Ammonia Water be added until of alkaline reaction, no turbidity or precipitation should take place, either before or after boiling, indicating the limit of Iron, Aluminium, Phosphates, etc, *U S P* A solution of the salt in Acetic Acid should not give a precipitate on saturation with Solution of Ammonia, *P G*

Barium Nitrate—The 1-50 solution obtained on boiling, by means of diluted Acetic Acid should not be at once affected by Barium Nitrate Solution, *P G*.

Silver Nitrate—The 1-50 aqueous solution obtained by dissolving in diluted Nitric Acid and boiling shall at the most be rendered only faintly opalescent after five minutes on the addition of Silver Nitrate Solution, *P G*

Potassium Ferrocyanide—The solution obtained by dissolving 1 gramme of the salt by the aid of Hydrochloric Acid in 50 c c of Water shall not become blue on the addition of 0.5 c c of Potassium Ferrocyanide Solution, *P G*

CRETA PRÆPARATA. *See p 456*

CALCII CHLORIDUM.

CALCIUM CHLORIDE

$\text{CaCl}_2, 2\text{H}_2\text{O}$, eq 145.85

FR, CHLORURE DE CALCIUM, GER, CALCIUMCHLORID

White, or nearly white, slightly translucent, hard fragments, having a sharp saline taste. Very deliquescent.

The official salt is obtained by the interaction of Hydrochloric Acid and Calcium Carbonate, the latter being added until the Acid is neutralised. The salt is dried at a temperature not exceeding 200°C (392°F). A slight dissociation and loss of Hydrochloric Acid occurs during the drying, and most commercial samples are alkaline.

The *USP* gives no specific temperature at which the salt should be dried, but states that when it has been overheated the solution has an alkaline reaction and a residue of Oxide remains undissolved, which, however, goes into solution in Hydrochloric Acid.

It should be kept in glass bottles with closely-fitting glass stoppers.

Solubility—1 in 1 of Water, 1 in 3 of Alcohol (90 p 63)

Medicinal Properties.—It increases the coagulability of the blood, and is therefore used in gastric, intestinal and pulmonary hæmorrhage, also in hæmophilia and aneurisms. Has been given internally also for chilblains, 20 grains night and morning, and in glandular enlargements, especially those of tubercular origin.

Given in pneumonia—*Pr* 1 263, *lin* 343

10 to 20 grains every 4 hours given in hæmophilia—*L*. '97, *n* 1061, *L*. '98, *n* 144, *B M J* '02, *i* 1141, *P J* '03, *i* 525

A small pledget of Wool soaked in a solution containing 30 grains to the oz of Water, used successfully in hæmophilia—*L*. '03, *i* 517

If pulmonary hæmorrhage persists, 2 or 3 drms. Chloride or Lactate of Calcium should be given in the course of a few hours—*T G* '07, 323

In pneumonia, 5 to 10 grains dissolved in Water every 4 hours, 1 minim of Elixir of Saccharin covers the taste of 10 grains, does not interfere with use of other remedies (Lauder Brunton)—*B M J* '07, *i* 616

Valuable in hæmorrhagic type of pneumonia, but its use deprecated in dry—*B M J* '07, *i* 1176

Can have no place in treatment of early stage of pneumonia (William Ewart).—*B M J* '07, *i* 779

In hæmorrhagic gastric oozing, 10 grains thrice daily (Hale White)—*L*. '06, *n* 1193.

Rapidly effective, in 3 or 4 grain doses thrice daily, in purpura in children (*B M J* '07, 1 199), and in 15 grain doses in boy of 16, p 1865

Given as a preventive in full doses for 3 days before any intranasal operation, and as a curative in 20 to 60 grain doses in hæmorrhage after extraction of teeth—*B M J* '07, 1 1054, 1 83

When used for hæmoptysis it must not be given for more than 3 or 4 days at a time, otherwise the blood will become less coagulable—*Pr* '07, 1 335

It markedly diminished the amount of albumen excreted by patients with Bright's disease of the non-polyuric form, but who passed 3 to 15 grammes albumen per litre—*L* '07, 1 841

The value of this salt in intestinal hæmorrhage given in 10 or 20 grain doses every 3 hours has been shown (*B M J* '04, 1453) 30 to 40 grains dissolved in a little Water and injected into the rectum, has also been recommended (*B M J* '04, 1 1635), objection has been taken to this method on account of disturbing patient (*B M J* '04, 1 1783), but it has been pointed out (*B M J* '05, 1 108) that good results in hæmoptysis have been accompanied by very little discomfort to patient

It is stated (*B M J* '06, 1 26) that this substance acts as a harmless and simple preventive against free bleeding when administered prior to an operation, and at the same time is not less efficient than other hæmostatic agents Administered in 1 drgm doses night and morning for 3 days previous to operation to prevent hæmorrhage in excision of the testicle in a case of tubercular disease of the seminal tract—*M P* '06, 1 38

Dose—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—*This Salt has a very unpleasant taste which is difficult to cover. Elixir Calcium Chloridi accomplishes this better than Liquorice*

Incompatibles—Lime salts and Potassium salts are mutually antagonistic physiologically—*B M J* '87, 1 1038

Official Preparations—Used in the preparation of *Æther Purus*

Not Official—*Elixir Calcium Chloridi, Liquor Calcium Chloridi, Syrupus Calcium Chloridi*

Foreign Pharmacopœias—Official in U.S., Hung (*Calcium Chloratum Fusum*) Port (*Chloreto de Calcio*), Fr (*Chlorure de Calcium Cristallisé* and *Chlorure de Calcium Fondu*), Mex (*Cloruro de Calcio*), Span (*Cloruro Calcio*) Not in the others

Tests—Calcium Chloride should answer the tests distinctive of Calcium given in the large type under *Calcium Carbonas Præcipitatus*. The aqueous solution yields with Silver Nitrate Solution a white curdy precipitate, insoluble in Nitric Acid, soluble in Ammonia Solution and in Potassium Cyanide Solution. A small quantity of the salt heated with Manganese Dioxide and Sulphuric Acid evolves a yellowish-green gas recognised as Chlorine by its odour and liberating Iodine from Potassium Iodide Solution, which on the addition of Starch Mucilage yields a blue coloration

The *B P* does not fix a limit for the amount of pure Calcium Chloride which the salt shall contain, the *U S P* requires that it shall contain not less than 99 p.c. of the pure salt. Neither Pharmacopœia describes a process of determination. The neutral aqueous solution may be titrated with Deci-normal Volumetric Silver Nitrate Solution, or the Calcium may be determined as Oxalate, the Oxalate washed, dried, ignited, and weighed as Calcium Oxide

The more generally occurring impurities are Aluminium, Iron, Magnesium, Carbonates and Hypochlorite. Aluminium, Iron and Magnesium, if present, may be detected by the Ammonia Solution test given below in small type under the heading of Ammonia,

Carbonates by an effervescence following the addition of Hydrochloric Acid HCl , by Ammonium Phosphate Solution after separation of the Calcium as Oxalate in presence of Ammonium Chloride and a trace of Ammonia Solution. Hypochlorite might have been a likely impurity in a salt made according to the process described in *BP* '85, but its presence in a sample prepared according to the *BP* '98 directions is unlikely, especially when dried at the temperature officially directed. The *USP* includes Phosphates as capable of detection by Ammonia Solution test, it also includes a time-limit test for Arsenic and Lead, with Hydrogen Sulphide and a test indicating a limit of Magnesium and alkalis, it is performed by completely precipitating the Calcium from 10 cc of a 1 in 20 aqueous solution by means of Ammonium Oxalate Test-solution, evaporating the filtrate to dryness and igniting, not more than 0.1 of a gramme of fixed residue should remain. The salt is not official in *P G*.

Ammonia—If to the aqueous solution of the salt Ammonia Water be added until of alkaline reaction, no turbidity or precipitation should take place, either before or after boiling, *USP*.

Hydrogen Sulphide—The aqueous solution of the salt (1-20) slightly acidulated with Hydrochloric Acid should not respond to the time-limit test for Arsenic or Lead, *USP*.

Not Official

LIQUOR CALCI CHLORIDI—Chloride of Calcium 1, Distilled Water 5—*BP* 1885

This has been incorporated in the *BP C* as follows—

Calcium Chloride 16, Distilled Water, *qs* to produce 100

ELIXIR CALCI CHLORIDI—Chloride of Calcium, 60 grains, Citric Acid, 20 grains, Aromatic Elixir (*USP*), to 1 fl oz

Syrupus Calcii Chloridi. *Syn* Elixir of Calcium Chloride—Calcium Chloride 12.50 Citric Acid 5, Distilled Water 15, Aromatic Syrup, *qs* to produce 100—*BP C*

CALX CHLORINATA. See p 298

Not Official

CALCI GLYCEROPHOSPHAS.

CALCIUM GLYCEROPHOSPHATE

FR, GLYCEROPHOSPHATE DE CALCIUM, GER, GLYCERINPHOSPHORSAURES CALCIUM, ITAL, GLICEROFOSFATO DI CALCIO, SPAN, GLICEROFOSFATO DE CAL

A white odorless crystalline powder

It is prepared by the action of Milk of Lime on Glycerophosphoric Acid, the product being purified by treatment with Alcohol

Solubility—1 in 22 of Water, less soluble in warm Water, and almost insoluble in boiling Water, insoluble in Alcohol (90 p c)

Medicinal Properties—It improves the general nutrition in neurasthenia

Dose—5 to 15 grains = 0.32 to 1 gramme dissolved in Water

Prescribing Notes—*Liquid Glycerophosphate preparations keep best when undiluted, and any dilution should be done when the medicine is taken, but it*

prescribed in a diluted form the diluent should be Chloroform Water, not Distilled Water

Foreign Pharmacopœias—Official in Belg., Fr., Mex (Glicerofosfato de Calcio), Span (Glicerofosfato de Cal), and Swiss (Calcium Glycerinophosphoricum)

Tests—Calcium Glycerophosphate dissolves in Water, yielding a solution which is neutral in reaction towards Litmus paper. The aqueous solution yields with Ammonium Oxalate a white precipitate soluble in diluted Hydrochloric Acid, when the cold aqueous solution is warmed a white precipitate is thrown out, the salt being less soluble in warm than in cold Water. A little of the salt heated in a tube blackens and evolves the characteristic irritating odour of Acrolein, when ignited on platinum foil it burns with a luminous flame, leaving a white residue, which dissolved in Nitric Acid yields a solution giving on the addition of Ammonium Molybdate Solution a yellow precipitate insoluble in Nitric Acid, soluble in Ammonia Solution.

The Glycerophosphates do not answer the usual reactions of the Phosphates, that is to say they do not give an immediate precipitate with Ammonium Molybdate Solution, nor do they immediately precipitate with Magnesium Ammonio-sulphate Solution. On boiling with a mineral acid the Glycerophosphoric Acid is decomposed, and the solution then responds to the tests for Phosphoric Acid. The aqueous solution of the salt yields a white precipitate with Lead Subacetate Solution. The percentage of Phosphoric Acid may be determined volumetrically by titration of the aqueous solution with Deci-normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator, and then titrating the solution with Deci-normal Volumetric Potassium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator. In the first instance one molecule of mineral acid corresponds to two molecules of Phosphoric Acid, and in the second two molecules of alkali correspond to one molecule of Phosphoric Anhydride.

The more generally occurring impurities are Arsenic, heavy metals, *eg*, Lead, Copper, and Iron, Chlorides, Phosphates, and Sulphates, readily carbonisable organic impurities, and free Glycerin.

Arsenic, if present, may be detected by Bettendorf's test, heavy metals by Hydrogen Sulphide in a solution rendered faintly acid with Hydrochloric Acid, Iron by Ammonium Hydrosulphide Solution, Chlorides by Silver Nitrate Solution in a solution rendered acid with Nitric Acid, Phosphates and Sulphates generally exist as insoluble salts and are readily detected by the solubility, and also by Barium Chloride or Nitrate Solution, organic impurities by charring when the salt is treated with concentrated Sulphuric Acid, and free Glycerin by treating the dry salt with Absolute Alcohol, filtering and evaporating off the solvent.

GLYCEROL GLYCEROPHOSPHATIS—Oudbear, 15 grains, Distilled Water, 10 fl oz. Boil for 10 minutes, filter and dissolve in the warm filtrate, Calcium Glycerophosphate, 160 grains, Potassium Glycerophosphate, 80 grains, Sodium Glycerophosphate, 80 grains, Magnesium Glycerophosphate, 80 grains, Iron Glycerophosphate (in scales), 40 grains, Citric Acid, 80 grains, then add Glycerin, 10 fl oz, Chloroform, 5 minims, Alcohol, 40 minims, Orange Flower Water (triple), 2 fl drms, Cherry Laurel Water, 3 fl drms, Distilled Water, *q s* to produce 20 fl oz.—*Bournemouth Formulary*

This has been incorporated in the *B P C* under the title **Glycerinum Glycerophosphatum Compositum** *Syn* Glycerol Glycerophosphatis

SYRUPUS GLYCEROPHOSPHATUM—The following is the formula given by Dr Robin, who introduced the preparation—Calcium Glycerophosphate, 6 grammes, Sodium, Potassium, and Magnesium Glycerophosphates, of each 2 grammes, Iron Glycerophosphate, 1 gramme, Tinctura Ignatæ Amara, 80 minims, Pepsin, 3 grammes, Maltine, 1 gramme, Tincture of Kola, 10 grammes, Syrup of Cherries, to 200 grammes.—*P J '95*, 1 1191

Nearly all commercial Syrups of the Glycerophosphates become turbid and throw down a bulky deposit on standing, and the same objection applies to **Syrupus Glycerophosphatum Compositus (B P C)**. The composition of

this latter preparation has been altered twice in about twelve months. It first of all contained Citric Acid, which was replaced by Acetic Acid, which is now replaced in the *B.P.C. Supplement* by Glycerophosphoric Acid.

FERRI GLYCEROPHOSPHAS (Iron Glycerophosphate) — Yellow or yellowish-green scales, or as a white powder, soluble in cold Water, more readily in hot. Introduced as a nervine tonic. Has been recommended in anæmia and especially chlorosis.

Dose — 5 to 10 grains = 0.32 to 0.65 gramme

Acidum Glycerophosphoricum — A clear, colourless, or pale yellowish-syrupy liquid. Sp. gr. 1.127. Chiefly used for the preparation of the salts, but rarely used medicinally.

CALCII HYDRAS.

CALCIUM HYDROXIDE

B.P. Syn. — SLAKED LIME

$\text{Ca}(\text{HO})_2$, eq. 73.47

A white odourless powder, having a caustic taste.

The Pharmacopœia directs that 'it should be recently prepared,' but this is unnecessary if air be excluded.

It should be kept in well-stoppered glass bottles, and protected as far as possible from contact with the air.

Solubility — Sparingly soluble in Water (1 in 900); the solution, on exposure to the air, soon acquires a film of Calcium Carbonate.

Medicinal Properties. — Antacid, astringent, sedative. The **Solution** (Lime Water) is useful in acid and gouty dyspepsia; in vomiting and diarrhœa of children, especially if given with the Milk, as it renders the curd less dense, in enteric fever it lessens the chances of hæmorrhage, also in the form of diluted **saccharated solution** to relieve chronic vomiting, and vomiting of pregnancy. The **Liniment** of Lime is applied to burns and scalds. When made with Linseed Oil it is known as Carron Oil.

Incompatibles — Vegetable and Mineral Acids, alkaline and metallic salts, Tartar Emetic.

Official Preparations — **Liquor Calcis** and **Liquor Calcis Saccharatus**. Used in the preparation of **Calcii Hypophosphis**, **Chloroformum**, **Extractum Ipecacuanhæ Liquidum**. **Lime Water** is used in the preparation of **Calcii Oxidum**, **Linimentum Calcis**, **Lotio Hydrargyri Flava** and **Lotio Hydrargyri Nigra**.

Not Official — Liniment for freckles, Carron Oil.

Foreign Pharmacopœias — Official in Fr. (**Hydroxide de Calcium**) and Span. (**Cal Apagada**). Not in the others.

Tests — Slaked Lime when dissolved in Water or in diluted Hydrochloric Acid and neutralised should afford the tests distinctive of Calcium given in the large type under **Precipitated Calcium Carbonate**. The aqueous solution should be alkaline in reaction to red Litmus paper and to Phenolphthalein Solution, and may be titrated with Normal Volumetric Hydrochloric or Sulphuric Acid Solution by the use of the latter indicator. 1 cc. of Normal Volumetric Acid Solution is equal to 0.036735 gramme.

of pure Calcium Hydroxide. When strongly heated the salt loses nearly one fourth of its weight of Water.

The more generally occurring impurities are Aluminium, Iron, Magnesium, Potassium, Sodium, Carbonates, Chlorides, Phosphates and Sulphates, and Silica if present is detected by the residue remaining insoluble in diluted Hydrochloric Acid, especially if, after first dissolving in Hydrochloric Acid the solution is evaporated to dryness and redissolved in Hydrochloric Acid, Aluminium, Iron and Phosphates are shown by the appearance of a precipitate when the Hydrochloric Acid Solution is rendered distinctly alkaline with Ammonia Solution, Magnesium by the precipitate produced with Ammonium Phosphate after separation of the Calcium as Oxalate in a solution containing Ammonium Chloride and a slight excess of Ammonia Solution. Potassium and Sodium may be detected, if present, in the residue after complete separation of the Calcium by Ammonium Oxalate. The absence of effervescence on the addition of Hydrochloric Acid is indicative of the absence of Carbonate, whilst Chloride may be detected in the Nitric Acid Solution of the salt by Silver Nitrate Solution and Sulphates in another portion of the same solution by means of Barium Chloride or Nitrate Solution. Phosphate may be detected by adding Ammonium Molybdate Solution to a solution of the salt containing some free Nitric Acid.

Preparations

LINIMENTUM CALCIS LINIMENT OF LIME

Solution of Lime, 1, Olive Oil, 1 (1 in 2)

Foreign Pharmacopœias—Official in Belg., Solution of Lime and Medicinal Oil, equal parts, Fr. (*Linim Calcaire*), Solution of Lime and Olive Oil, equal parts, Ital., Lime Water and Olive Oil, equal parts, Jap. and Mex., Lime Water 1, Sesame Oil 1, Port., Lime Water 9, Oil of Almonds 1, Span., (*Linimento Oleo-Calcareo*) Lime Water 65, Oil of Almonds 35, Austr., Dan. (*Linimentum Calcicum*), Dutch, Russ., Swed., Swiss and U.S., Solution of Lime and Linseed Oil, equal parts. All by weight, except U.S. Not in the others.

LIQUOR CALCIS SOLUTION OF LIME *B.P. Syn*—LIME WATER

A saturated Solution of Calcium Hydroxide (washed free of Chlorides) in Distilled Water. It should contain about $\frac{1}{2}$ grain in the fl. oz.

It should be kept in well-stoppered glass bottles of a dark amber colour, and preserved as far as possible from contact with the air. It is best kept in full bottles.

When freshly prepared it forms a colourless and odourless liquid possessing a peculiar alkaline taste. It gradually absorbs Carbonic Anhydride from the air, and a crust of Calcium Carbonate forms on the surface of the liquid.

The Hydrate is less soluble in hot than in cold Water, and, if saturated, Lime Water should precipitate on boiling.

Dose.—1 to 4 fl. oz. = 28 4 to 113·6 c.c.

It is more palatable when given in Milk

So-called aerated 'Lime Water' is sold in syphons, but we understand that it is aerated with Carbonic Acid gas, and in that case the name is misleading

Foreign Pharmacopœias—Official in Austr and Hung (Aqua Calcis), Dan, Dutch, Norw and Swed (Solutio Hydratis Calcis), Fr (Eau de Chaux), Ger and Jap (Aqua Calcariae), Ital (Acqua di Calce), Mex, Port and Span (Agua de Cal), Russ (Calcium Caustica Soluta), Belg, Swiss (Calcium Hydricum Solutum), U S, Liquor Calcis

Water becomes saturated with much less Lime than ordered in any of the Pharmacopœias, therefore Liquor Calcis is of the same strength in all

Tests.—Lime Water is strongly alkaline in reaction towards Litmus. It should yield the tests distinctive of Calcium described in the large type under Calcium Carbonate. It is officially required to contain 0.153 p.c. w/v of pure Calcium Hydroxide, as volumetrically determined by titration with Deci-normal Volumetric Sulphuric Acid Solution. Neither the *BP* nor the *PG* specify a particular indicator, the *USP* mentions Phenolphthalein Solution. The *PG* requires not less than 0.148 p.c. w/v, nor more than 0.166 p.c. w/v, the *USP* not less than 0.14 p.c. w/v, of pure Calcium Hydroxide.

The liquor should be free from the more generally occurring impurities mentioned under Calcium Hydrate, and in addition should be free from objectionable traces of Lead.

LIQUOR CALCIS SACCHARATUS. SACCHARATED SOLUTION OF LIME

Calcium Hydroxide, 1, Refined Sugar, in powder, 2, Distilled Water, 20 (about 1 in 62)

Contains about 8 grains of CaO in 1 fl oz. 1 oz. = about 14 fl oz. Lime Water

As suggested in our former editions the Sugar should be first dissolved in the Water, and the Calcium Hydroxide added to the Solution, after a few hours' occasional agitation, decant.

It should be kept in well-stoppered glass bottles of a dark amber colour, which should be kept full, and preserved as far as possible from contact with the air.

The Hydrate in this case also is less soluble in the hot than in the cold, and the liquor precipitates on boiling, but clears again on cooling.

A clear, colourless, and odourless liquid possessing a peculiar sweetish and alkaline taste.

Dose.—20 to 60 minims = 1.2 to 3.6 c.c.

Foreign Pharmacopœias—Official in Hung (Aqua Calcis Saccharata). Not in the others.

Tests. *Saccharatus* of Lime has a specific gravity of about 1.055, and is officially required to indicate 2.31 p.c. w/v of pure Calcium Hydroxide, as volumetrically determined by titration with Normal Volumetric Sulphuric Acid Solution. In this case also the *BP* specifies no indicator of neutrality. Phenolphthalein Solution is best for the purpose.

The liquor should be free from the impurities usually associated with the Hydroxide from which it is prepared, and in addition should not contain objectionable traces of Lead

Not Official

LINIMENT FOR FRECKLES—Liniment of Lime, 8, Solution of Ammonia, 1, mix

CARRON OIL—Equal parts of Linsced Oil and Lime Water, shaken to form a cream

One of the best applications to burns or scalds, more particularly when 1 or 2 p c of Phenol has been added

This has been incorporated in the *B P C* under the title *Linimentum Calcis cum Oleo Lini*

CALCII HYPOPHOSPHIS.

CALCIUM HYPOPHOSPHITE

FR, HYPOPHOSPHITE DE CALCIUM, GER, CALCIUM HYPOPHOSPHORUM, ITAL, IPOFOSFITO DI CALCIO, SPAN, HIPOFOSFITO CALCICO

$\text{Ca}(\text{PH}_2\text{O}_2)_2$, eq 168.83

A white, odourless, lustrous crystalline salt, or as a white crystalline powder, having a nauseous bitter taste

It should be kept in well-stoppered bottles and in a cool place. The salt readily undergoes oxidation, and on this account great care should be exercised in mixing it with substances which readily part with Oxygen, *e g*, Chlorates, Nitrates, and Peroxides

It may be obtained by the combined action of Calcium Hydroxide, Phosphorus, and Water, the salt being purified by recrystallisation

Solubility—1 in 8 of Water, and scarcely more soluble in boiling Water. Insoluble in Alcohol (90 p c)

Medicinal Properties.—Similar to those of Phosphorus, but without its unpleasant effects. Given in cases of nervous and general debility, it is by some supposed to be useful in phthisis

No satisfactory evidence that Hypophosphites can influence nutrition in any way differently from other indifferent salts, and the whole of the Hypophosphite administered can be recovered unchanged from the urine—*B M J* '06, ii 1460

Dose—3 to 10 grains = 0.2 to 0.65 gramme

Prescribing Notes—Usually given in mixtures or in one of the various forms of Syrup

Not Official—Glycerola Hypophosphitum, Syrupus Hypophosphitum, Syrupus Calci Hypophosphitis (*Squire*) and Syrupus Calci et Sodii Hypophosphitum

Foreign Pharmacopœias—Official in Belg, Jap and Swiss (Calcium Hypophosphorosum), Dutch and Norw (Hypophosphis Calcicus), Fr (Hypophosphite de Chaux), Ital (Ipofofrito di Calcio), Mex (Ipofofrito de Calcio), Port (Hypophosphito de Cal), Span (Ipofofrito Calcico), U S Not in the others

Tests—The distinguishing tests for Calcium Hypophosphite are

(1) that when heated to redness the crystals evolve spontaneously inflammable Hydrogen Phosphide and Hydrogen, and leave a reddish-coloured residue, (2) an aqueous solution rapidly reduces Potassium Permanganate Solution, yielding if the proportionate quantity of Permanganate be employed a practically colourless filtrate, (3) the aqueous solution reduces Mercuric Chloride Solution first to Mercurous Chloride and ultimately more completely to metallic Mercury. Upon its property of reducing Potassium Permanganate Solution is founded the official quantitative test of its purity, which requires that when 0.25 gramme of the salt is boiled for 10 minutes with 0.6 gramme of Potassium Permanganate in solution, and the liquid filtered, the filtrate shall be practically colourless. Several methods have been proposed with a view to affording a more accurate quantitative determination. Tyrer has suggested (*P. J.* '97, ii 150) reduction of Copper Solution, previously eliminating any impurity likely to affect the result, by treatment with Barium Chloride Solution. Jowett points out (*Y. B. P.* '98, 412, *C. D.* '98, ii 300) that Barium Phosphite is slightly soluble in Water, which would affect the results obtained, and proposes the following method—About 0.3 gramme of the dried salt is dissolved in 10 c.c. of Water, 3 c.c. of a 10 p.c. Lead Acetate solution added, and the mixture allowed to stand 12 hours. It is then filtered, the precipitate thoroughly washed, and the washings added to the filtrate, which is acidified with Hydrochloric Acid, and then saturated with Hydrogen Sulphide, boiled, filtered, and the Lead Sulphide thoroughly washed. The mixed washings and filtrate are then evaporated to a low bulk and 5 c.c. Hydrochloric Acid and 1 gramme Potassium Chlorate added, and gently heated for half an hour, then concentrated to about 20 c.c., and the Phosphate finally determined either gravimetrically or volumetrically by the usual methods. The *U. S. P.*, although requiring that it shall contain not less than 98 p.c. of pure Calcium H₂P₂O₇, gives no process for its determination. The salt is not official in the *P. G.*

The more generally occurring impurities are Aluminium, Arsenic, Copper, Iron, Lead, Magnesium, Potassium, Sodium, Chlorides or Sulphates, Phosphates or Phosphites. The *B. P.* groups these collectively. A 1 in 20 aqueous solution of the salt acidified with Hydrochloric Acid should yield no turbidity or darkening in colour on the addition of Hydrogen Sulphide indicating the absence of Arsenic, Copper and Lead, nor on the subsequent addition of Ammonia Solution should any appreciable darkening in colour ensue, indicating the absence of more than a trace of Iron. The 1 in 20 aqueous solution, after the complete precipitation of the Calcium as Oxalate, should afford no turbidity on the addition of Sodium Phosphate Solution, indicating the absence of Magnesium. The *U. S. P.* does not include tests for Aluminium, Magnesium, Potassium, Sodium or Phosphites; the modified Gutzeit's test is employed in testing for Arsenic, and the time-limit test for heavy metals. The *B. P.* employs Lead Acetate Solution to detect Phosphates and Phosphites, but no commercial sample has been found which does not give more or less precipitate or turbidity with Lead Acetate, which also precipitates Sulphates and Sulphites.

The *USP* takes the solubility of the salt in Water as a criterion of the absence of Phosphate and Sulphate

Modified Gutzzeit's Test—If 5 c c of an aqueous solution of the salt (1-10) be measured into a beaker containing 3 c c of Nitric Acid diluted with about 10 c c of Water and evaporated to dryness on a water bath the residue should not respond to the modified Gutzzeit's test for Arsenic, *USP*

Time-limit Test—A (1-20) aqueous solution of the salt acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*

Not Official

GLYCEROLA HYPOPHOSPHITUM (*Squire*)—Calcium, Potassium, and Sodium Hypophosphites, of each 1, dissolve these in Water, 40, filter and add Sugar, 40, Orange flower Water, 2, Cherry laurel Water, 2, dissolve and add Glycerin, 12, and filter

Dose—1 to 2 fl drm = 3 6 to 7 1 c c

Glycerol Hypophosphitis—Hypophosphite of Potassium, 160 grains, Hypophosphite of Calcium, 160 grains, Hypophosphite of Manganese, 80 grains, Hypophosphite of Quinine, 80 grains, Hypophosphite of Strychnine, 2½ grains, Strong Solution of Hypophosphite of Iron (*BPC*), 4 fl oz, Hypophosphorous Acid, 2 fl drm, Distilled Water, 3 fl oz, Glycerin, to produce 20 fl oz — *Bournemouth Formulary*

This has been incorporated in the *BPC* as follows —

Glycerinum Hypophosphitum *Syn* GLYCEROL HYPOPHOSPHITIS — Calcium Hypophosphite, 1 50, Manganese Hypophosphite, 0 75, Potassium Hypophosphite, 1 50, Quinine Hypophosphite, 0 75, Strychnine Hypophosphite, 0 025, Strong Solution of Ferric Hypophosphite, 20, Hypophosphorous Acid, 10, Distilled Water, 15, Glycerin, q s to produce 100 — *BPC*

SYRUPUS HYPOPHOSPHITUM — Calcium Hypophosphite, 45 grammes, Potassium Hypophosphite, 15 grammes, Sodium Hypophosphite, 15 grammes, Diluted Hypophosphorous Acid, 2 grammes, Sugar, 650 grammes, Tincture of Fresh Lemon Peel, 5 c c, Water, q s. to make 1000 c c — *US*

SYRUPUS CALCII HYPOPHOSPHITIS (*Squire*)—Dissolve Calcium Hypophosphite, 4 in Distilled Water, 38 add Sugar, 59, dissolve without heat and filter the Syrup

Dose—1 fl drm = 3 6 c c containing 3 grains = 0 2 gramme

The *BPC* Syrup is only one third the strength, necessitating an excessive quantity of Syrup for a full dose of the salt

SYRUPUS CALCII ET SODII HYPOPHOSPHITUM (*USNF*) — Calcium Hypophosphite, 35, Sodium Hypophosphite, 35, Hypophosphorous Acid (*USP*), 1 5, Sugar, 775, Water, sufficient to make 1000

1 fl drm contains 2 grains each of Calcium Hypophosphite and Sodium Hypophosphite

The *BPC* has incorporated the old formula (*USNF* 1896) which employs Citric Acid

Not Official CALCII IODAS

CALCIUM IODATE

A white, or more usually yellowish, very deliquescent, crystalline powder antiseptic and disinfectant Under the name Calcinol it has been used as an antiseptic dressing in place of Iodoform — *L* '00, 11 1887, *PJ* '01, 1 27

Dose—1 to 3 grains = 0 06 to 0 2 gramme, two or three times daily in Water, as an intestinal antiseptic

Tests.—The aqueous solution affords the tests characteristic of Calcium given in large part under Calcium Carbonate

If a few drops of Silver Nitrate Solution a white precipitate, sparingly soluble in Water and in dilute Nitric Acid, but readily soluble in Ammonia Solution. When mixed with Potassium Iodide Solution it yields on the addition of Tartaric Acid Solution or a solution of an acid Sulphate and Starch Mucilage a fine blue coloration. Barium Chloride Solution produces a white precipitate practically insoluble in Water and difficultly soluble in Nitric Acid. On the addition of Sulphurous Acid to an aqueous solution Iodine is liberated, and may be recognised by the violet colour of its solution in Carbon Bisulphide or by the blue colour produced on the addition of Starch Mucilage

Not Official.

CALCII IODIDUM.

This salt may be obtained from concentrated solutions in crystalline needles. By evaporating its solution to dryness it may be obtained as a white deliquescent fusible mass which crystallises on cooling. It is readily soluble in Water and in dilute Alcohol. It has recently (*B M J* '06, i, 138) been shown that great benefit is derived from its use in the treatment of ulcers, it had a remarkable effect in reducing thick callous edges into thin healing ones. It is given in doses of 2 to 4 grains (0.13 to 0.26 gramme)

Of great value in leg ulcers.—*B M J* '07, i, 991

Suggested that the gastritis which has sometimes followed its administration was due to free Iodine given off by the salt on exposure to a bright light or air. It should, therefore, be kept unexposed to light, and dispensed in suitably coloured bottles.—*B M J*, '07, i, 1464

CALCII PHOSPHAS.

CALCIUM PHOSPHATE.

FR, PHOSPHATE DE CALCIUM, GER, CALCIUMPHOSPHAT, ITAL, FOSFATO DI CALCIO, SPAN, FOSFATO CALCICO

A light, white, odourless and tasteless amorphous powder

It is generally obtained commercially from Bone Ash. The Bone Ash is treated with Hydrochloric Acid and the resulting solution is precipitated with dilute Ammonia Solution, the precipitate being washed with cold Water to free it from Ammonium salts and dried at a temperature not exceeding 100° C (212° F). It is officially mentioned that it may be prepared by the interaction of Calcium Chloride and Sodium Phosphate, but it has been stated (*C D* '02, i, 190) that the salt produced by precipitation with the official Di-Sodium Hydrogen Phosphate is Di-Calcium Hydrogen Phosphate, and only in the event of Tri-Sodium Phosphate being used as a precipitant is a salt corresponding to the official *Calcii Phosphas* produced. A specimen produced by precipitation of Calcium Hydrogen Phosphate when dissolved in Hydrochloric Acid and reprecipitated with Ammonia Solution showed a loss of over 29 p c, whereas the official requirement is not less than 5 p c.

Solubility—Insoluble in Water, soluble in Dilute Hydrochloric Acid or Diluted Nitric Acid.

Medicinal Properties—For rickets and mollities ossium, and other conditions of malnutrition, said to be useful in scrofulous affections, to promote union of bone fractures, in tardy teething, and in anæmia, given to counteract the draining of Phosphates during pregnancy and lactation, and to prevent decay of the teeth and toothache during pregnancy

Dose—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—*More commonly ordered in smaller doses. Given as a powder, or in the form of Syrup*

Official Preparations—Contained in *Extractum Euonymi Siccum* and *Pulvis Antimonialis*

Foreign Pharmacopœias—Official in Austr, Belg, Ger, Hung, Russ and Swiss (*Calcium Phosphoricum*), Dutch and Swed (*Phosphas Calcicus*), Dan (*Phosphas Calcicus Præcipitatus*), Fr (*Phosphate Diacide de Calcium*, *Phosphate Monoacide de Calcium* and *Phosphate Neutre de Calcium*), Ital (*Fosfato Bicalcico*), Jap (*Calcium Phosphoricum Præcipitatum*), Mex (*Fosfato de Calcio*), Port (*Phosphato de Cal*), Span (*Fosfato DiCalcico* and *Fosfato TriCalcico*), US (*Calcii Phosphas Præcipitatus*) Not in Norw

Tests—Calcium Phosphate when dissolved in Hydrochloric Acid answers the tests distinctive of Calcium given in the large type under Calcium Carbonate. Its solution in diluted Nitric Acid gives with Ammonium Molybdate Solution a yellow precipitate, which, when filtered off and washed, dissolves in Ammonia Solution, and affords on the addition of Magnesium Ammonio-sulphate Solution a white crystalline precipitate. The *BP* includes a method of gravimetrically determining the purity of the salt requiring that its Hydrochloric Acid Solution shall yield a white precipitate amounting to 95 p.c. by weight when reprecipitated with diluted Ammonia Solution, washed with cold Water and dried at 100° C (212° F). The *USP* states that it shall contain not less than 99 p.c. of pure Calcium Phosphate, but gives no method for its determination. The *PG* neither gives the requisite percentage nor a method of determination. The salt official in the *PG* is the Di-Calcium Phosphate and not the Tri-Calcium Phosphate.

The more generally occurring impurities are Aluminium, Arsenic, Copper, Iron, Lead, Magnesium, Carbonates, Chlorides, Silica and Calcium Oxalate. With the exception of the last named the *BP* group these collectively without any regard for their relative importance. Arsenic, the most important, is examined for in the *USP* by the modified Gutzeit's test, in the *PG* by Bettendorf's test. The *USP* employs the time limit test, and *PG* Hydrogen Sulphide for the detection of Copper and Lead. Neither the *USP* nor *PG* includes a test for Magnesium or Silica. Calcium Oxalate is an unlikely impurity, and it has been questioned (*CD '02*, 1190) whether the *BP* test for 'absence of Calcium Oxalate' is not a slip for 'absence of Alumina'. Both *USP* and *PG* state that when moistened with Silver Nitrate Solution the salt is coloured yellow, the *USP* mentions 'either before or after ignition,' which

distinguishes it from Acid Calcium Phosphate, which, after ignition, remains white, when so moistened

Potassium Sulphate—5 c c of the solution prepared by means of Nitric Acid strongly acidified with Nitric Acid should yield with 1 c c of Potassium Sulphate T S no turbidity upon standing, indicating the absence of Barium, *U S P*

Silver Nitrate—5 c c of a 1-20 solution prepared by the aid of Nitric Acid and acidulated with Nitric Acid yields with 0.5 c c of Silver Nitrate T S not more than a slight turbidity, *U S P*. The *P G* requires that a (1-20) aqueous solution of the salt obtained with Nitric Acid should be rendered at most opalescent with Silver Nitrate T S after 2 minutes

Barium Nitrate—If 1 part of Calcium Phosphate be shaken with 2 parts of Water, and the mixture filtered, the filtrate, after addition of Acetic Acid, should not be affected by Barium Nitrate Solution, *P G*

Stannous Chloride—A mixture of 1 gramme Calcium Phosphate and 8 c c of Stannous Chloride Solution should not assume a dark colour in the course of an hour, *P G*

Hydrogen Sulphide—The aqueous solution (1-20) obtained with Nitric Acid should give with excess of Solution of Ammonia on the addition of Hydrogen Sulphide T S a pure white precipitate, *P G*

Gutzzeit's Test—5 c c of a 1 in 10 solution of Calcium Phosphate in diluted Hydrochloric Acid should not respond to the modified Gutzzeit's test for Arsenic, *U S P*

Time-limit Test—An aqueous solution (1-20) obtained by shaking the salt with Water, adding Hydrochloric Acid drop by drop and heating until solution is effected, should not respond to the time-limit test for heavy metals, omitting the addition of Ammonia Solution, *U S P*.

Not Official.

CALCII SULPHAS.

CALCIUM SULPHATE.

SULPHATE OF LIME CALCINED GYPSUM PLASTER OF PARIS

Fr, SULFATE DE CALCIUM, GER, GYPSBRANNTER GIPS, ITAL, GESSO CALCINATO, SPAN, Yesso

Native Calcium Sulphate (CaSO_4 , $2\text{H}_2\text{O}$, eq 170.81) rendered nearly anhydrous by heat

A white odourless and tasteless powder

It should be kept in well-closed jars, and should be protected as far as possible from moisture

Foreign Pharmacopœias.—*U S P*, *B P*, *Pharmacopœia Germanica*, Ger, Hung, Jap, Mex, Norw, Russ, Swed, Sw, *Pharmacopœia Scandinavica*, *Pharmacopœia Sibirica*

The native salt is used for the preparation of Calc Sulphurata

Tests—The *U S P* test for Calcium Sulphate is that when mixed with half its weight of Water it should form a smooth paste, rapidly becoming hard. The *P G* states that when so mixed it shall harden within 5 minutes. The saturated aqueous solution answers the tests characteristic of Calcium given in the large type under Calcium Carbonate, it should be neutral to Litmus paper, and should on the addition of Barium Chloride Solution yield a white precipitate, insoluble in Hydrochloric Acid

The more generally occurring impurity is Calcium Carbonate and the salt should not yield any effervescence when added to dilute Hydrochloric or Nitric Acid.

ALX. SULPHURATA (Calcii Sulphuratum) See p. 302,

Not Official CALENDULA

COMMON MARIGOLD

The Florets of *Calendula officinalis*, L

Foreign Pharmacopœias —Official in U S Not in the others

TINCTURA CALENDULÆ FLORUM —1 of Marigold Flowers, dried, in No 20 powder, percolated with Alcohol (60 p c), to yield 5

This is included in the *B P C Formulary* 1901

Calendula, in No 20 powder, 20, Alcohol (95 p c), 100 Prepared by percolation —*U S P*

This has been incorporated in the *B P C*

Medicinal Properties —Used as an application for sprains and bruises, internally for amenorrhœa

Dose —5 to 20 minims = 0 3 to 1 2 c c

Foreign Pharmacopœias —Official in U S, 1 in 5 (Alcohol 94 9 p c) Not in the others

Not Official CALOTROPIS

Syn —MUDAR

The dried root bark of *Calotropis procera*, R Br, and of *Calotropis gigantea*, R Br, freed from the outer corky layer, dose 3 to 10 grains = 0 2 to 0 65 gramme, as a tonic, as an emetic 30 to 60 grains = 2 to 4 grammes, and **Tinctura Calotropis** (1 in 10), dose 30 to 60 minims = 1 8 to 3 6 c c, are official in the *Ind* and *Col Add* for India and the Eastern Colonies

CALUMBÆ RADIX.

CALUMBA ROOT

FR, RACINE DE COLOMBO, GER, KOLOMBOWURZEL, ITAL, COLOMBO,
SPAN, RAIZ DE COLOMBO

The Root of *Jateorhiza Calumba*, Miels, cut in nearly circular transverse discs and dried

The dried stem of *Coscinum fenestratum*, Colebr, (false Calumba Root) is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Medicinal Properties —A bitter stomachic useful in chronic atonic dyspepsia, in promoting appetite, stimulating the gastric functions and removing flatulence Given in convalescence from acute diseases, combined with alkalis or Bismuth Like other bitters, Calumba ought not to be given in gastric ulcer, in acute gastritis, or when there is pain, nor ought it to be given for too long

Prescribing Notes —Given in the form of *Infusum*, *Liquor Concentratus*, or *Tincture* with other medicines It is one of the few bitters that can be given with salts of Iron

Official Preparations —*Infusum Calumbæ*, *Liquor Calumbæ Concentratus*, and *Tinctura Calumbæ*.

Not Official—*Extractum Calumbæ, Fluidextractum Calumbæ, Infusum Calumbæ Concentratum*

Foreign Pharmacopœias—Official in all

Descriptive Notes—*Calumba* root as met with in commerce varies much in character and quality, some specimens being of fresh appearance and yellowish tint, and others dull or greyish-yellow in colour and badly dried, and sometimes worm-eaten. The transverse or oblique slices average about $1\frac{1}{2}$ inch (37 mm) in diameter and about $\frac{1}{2}$ inch (8 mm) in thickness. The *BP* defines the size as about $\frac{1}{2}$ to $\frac{3}{4}$ inch (3 to 12 mm) or more in thickness, and about 1 to 2 inches ($2\frac{1}{2}$ to 5 cm) or more in diameter. The bark is $\frac{1}{4}$ inch or more in thickness, and harder and less shrunken than the starchy parenchymatous central portion, which is more or less depressed. The whole surface has a number of slender lines radiating from the centre and the bark is marked off by a darker cambium ring. The taste is bitter and mucilaginous and the odour characteristic. Under the microscope the distinctive features are the large irregular starch grains (0.09 mm *P G*), a stellate hilum, and the irregularly thickened yellow sclerenchymatous cells in the bark, containing rhomboidal crystals of Calcium Oxalate. The active principle is not Berberine, as formerly supposed, but consists of two alkaloids resembling it (*P J* (4) xvi p 341). The infusion made with cold Water to avoid dissolving starch should be strained and then boiled to destroy a ferment or oxydase which causes the infusion, at first neutral, to become acid and turbid. To prevent the gum present from readily dissolving, the root is usually cut into small pieces instead of being pounded.

The root should be selected of good quality and of moderate size, as the larger pieces have less bark in proportion, and it is in this part that the activity of the drug chiefly resides. If old it is usually darker and duller in colour and more or less worm-eaten. A very bright colour usually indicates that the root has been washed and dried and may have lost some of its active principle. *Calumba* root is not often adulterated, sometimes small pieces with prominent woody wedges are occasionally found mixed with *Calumba*, but do not differ in the character of the starch or in taste, and are apparently pieces of the short underground stem from which the fusiform roots proceed. Another root has recently been found mixed with *Calumba*, resembling it in shape and size, but with a reddish tint, distinctly radiate structure, and containing bundles of acicular raphides in some parenchymatous cells and sphaeraphides in others.

Preparations

INFUSUM CALUMBÆ. INFUSION OF CALUMBA

Calumba Root, 1; Distilled Water, cold, 20, infuse for half an hour, strain (1 in 20)

Cold Water is used to avoid solution of the Starch which exists in the root,

Dose.— $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 c.c.

A corresponding preparation, **Infusum Coccinii**, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Foreign Pharmacopœias—Official in Belg, 1 of Fluid Extract in 20, Ital, 1 in 20, Span, 1 in 100 Not in the others

LIQUOR CALUMBÆ CONCENTRATUS. CONCENTRATED SOLUTION OF CALUMBA

An aqueous preparation of Calumba (preserved by the addition of Alcohol) 1 of Calumba in 2 of Liquor

It contains 22.5 p c by volume of Alcohol (90 p c)

Dose — $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 c c

Tests—Concentrated solution of Calumba has a specific gravity of 0.990 to 0.996, it contains about 4 p c w/v of total solids and about 20 p c w/v of Absolute Alcohol

A corresponding preparation, **Liquor Coccinii Concentratus**, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

TINCTURA CALUMBÆ TINCTURE OF CALUMBA

1 of Calumba Root in No 20 powder, macerated with 10 of Alcohol (60 p c) (1 in 10)

Dose — $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c c

Tests—Tincture of Calumba has a specific gravity of 0.915 to 0.920, it contains about 1 p c w/v of total solids and about 60 p c w/v of Absolute Alcohol

A corresponding preparation, **Tinctura Coccinii**, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Foreign Pharmacopœias—Official in Belg, Fr, Mex, Port, Span, Swed, Swiss and U S, 1 in 5, Ital and Jap, 1 in 10, all except U S by weight Not in the others

Not Official

EXTRACTUM CALUMBÆ—Calumba Root exhausted with Alcohol (60 p c) and the product evaporated to a pill consistence 16 parts of Root yield 1 to 1 $\frac{1}{2}$ parts of Extract

This has been incorporated in the *B P C*

Dose — $\frac{1}{2}$ to 2 grains = 0.03 to 0.13 gramme

Foreign Pharmacopœias—Official in Austr, Dutch, Hung and Span, made with Alcohol (70 p c), Belg, Ital and Mex, made with Alcohol (60 p c), Port, made with Alcohol (65 p c), Jap, made with Alcohol (45 p c), U S, **Fluid Extract** only, made with Alcohol (about 66 p c) Not in Dan, Fr, Ger, Norw, Russ or Swed

FLUIDEXTRACTUM CALUMBÆ (U S P)—A 1 in 1 Fluid extract prepared by macerating and percolating 1 of the Root with a mixture of 7 of Alcohol (94.9 p c) and 3 of Water. The Fluid Extract of the *Belg Pharm* is prepared from the Root with Alcohol (30°), the resultant fluid extract being required to contain at least 13 p c of dry residue

INFUSUM CALUMBÆ CONCENTRATUM—Calumba Root, in No 10 powder, 40, Alcohol (90 p c), 25, Dilute Chloroform Water (1 in 100), *q s* to make 100. Prepare by macero expression, the reserved liquid should be heated to 85° C, for 5 minutes before adding the Alcohol. **Dose** — $\frac{1}{2}$ to 1 fl drim—*Farr and Wright, P J '06, 1 165, '07, 1 621, C D '06, 1 252, Y B P '07, 247*
This appears in the *B P C*

CALX.

LIME

FR, OXYDE DE CALCIUM, GER, GEBRANNTER KALK, ITAL, OSSIDO DI CALCIO;
SPAN, CAL VIVA

Calcium Oxide, CaO , eq 55 59, obtained by calcining Chalk, Limestone, or Marble

White or whitish, hard, odourless masses, possessing a characteristic caustic taste When exposed to the atmosphere Lime rapidly absorbs Water and Carbonic Anhydride, and should therefore be kept in well-closed vessels, and protected as far as possible from contact with the air

Solubility.—Decomposed by Water, forming Calcium Hydrate, under which heading the solubility is given

Foreign Pharmacopœias—Official in all

Tests—The distinctive test for Lime is the heat evolved when mixed with about half its weight of Water, the masses swelling up and crumbling to powder The *BP* mentions rather less than its own weight of Water, the *PG* with half its weight of Water, and the *USP* with about half its weight of Water, the *USP* and the *PG* stating, in addition, that with from 3 to 4 parts of Water, it forms a smooth magma When mixed with Water and dissolved by the aid of Hydrochloric Acid, it yields a solution conforming to the distinctive tests for Calcium given in the large type under *Calci Carbonas Præcipitatus* The aqueous solution is alkaline in reaction towards red Litmus and to Phenolphthalein Solution The *USP* requires that in the anhydrous state it shall contain not less than 90 p c of pure Calcium Oxide, the *BP* and *PG* specify no definite percentage Neither Pharmacopœia refers to any method of quantitative determination

The more generally occurring impurities are Aluminium, Iron, Magnesium, Potassium, Sodium, Carbonates, Chlorides, Phosphates, Sulphates, and Silica. These are grouped collectively in the *BP* The methods adopted for their detection in the case of Calcium Hydrate are applicable here also The *PG* includes only a test for Carbonates and residue insoluble in Nitric Acid, the *USP* a limit of Carbonate, and a limit (0.5 p c) of matter insoluble in Hydrochloric Acid

CALCI HYDRAS.—See p 286

CALX CHLORINATA.

CHLORINATED LIME

FR., CHLORURE DE CHAUX, GER., CHLORKALK, ITAL, CLORURO DI CALCE.

A white, or greenish-white, dry, amorphous powder, evolving a strong chlorinnaceous odour Is a mixture of variable composition, consisting chiefly of Calcium Hypochlorite

As it becomes moist and gradually decomposes on exposure to the air, it should be preserved in well closed vessels in a cool and dry place

Solubility—Partially soluble in Water and in Alcohol (90 p c) Decomposed by acids with formation of Hypochlorous Acid, which in the case of Hydrochloric Acid reacts with it to form Chlorine

Medicinal Properties—Chiefly used as a disinfectant A solution of 1 of Calx Chlorinata to 120 of Water is very powerfully antiseptic and is neither toxic nor caustic Acts best at a temperature of 110° to 120° F

A remarkably efficacious and absolutely harmless antiseptic Useful as an external application in surgical, ophthalmic and gynaecological practice and also as an injection in affections of the rectum and bladder See also below, Liquor Calcis Chlorinatae

Bleaching Powder frequently consists of little else than inert Calcium Chloride and Carbonate, the active Chlorine being spent Generally speaking, it only destroyed the least resistant microbes, though in the case of anthrax spores on linen this substance proved more effective than Carbolic Acid—*London County Council's Report on Disinfectants, L '02, i 759*

Official Preparations—Liquor Calcis Chlorinatae Used in the preparation of Chloroform and Liquor Sodæ Chlorinatae

Not Official—Liquor Potassæ Chlorinatae

Foreign Pharmacopœias—Official in Dan, Norw, Swed (Calx Chlorata), Austri and Russ (Calcium Hypochlorosum), Fr (Chlorure de Chaux), Belg, Ger, Hung, Jap and Swiss (Calcaria Chlorata), Ital (Cloruro di Calce), Mex (Hipoclorito de Calcio Impuro), Port (Cal Chlorada), Span (Hipoclorito Calcico Clorinado), US (Calx Chlorinata), Hung and Norw, contain 20 p c of available Chlorine, Austr, Belg, Dan, Ger, Jap, Russ, Swed and Swiss, 25 p c, Ital, 28.6 p c, Fr 35 p c, Span, 32 p c, US not less than 30 p c, Port, not indicated Not in Dutch

Tests—The distinguishing tests for Calx Chlorinata are that when treated with Hydrochloric Acid it evolves a yellowish gas which at first reddens and then bleaches moistened blue Litmus paper, its aqueous solution decolorises Indigo Sulphate Solution, its Acetic Acid solution yields with Ammonium Oxalate Solution a white precipitate, insoluble in Acetic Acid, soluble in Hydrochloric Acid It is officially required to contain 33 p c (more correctly 32.93 p c) of available Chlorine as indicated by titration of the Iodine liberated on mixing 0.5 gramme of the salt with 1.5 grammes of Potassium Iodide dissolved in 200 c c of Water and 6 c c of Hydrochloric Acid, at least 46.8 c c of Volumetric Sodium Thiosulphate Solution should be required A detailed comparison of the *U.S.P.* and *P.G.* processes will be found below in the small type under the heading of Volumetric Determination It should be noted that only a good and well-kept sample will yield the percentage of Chlorine required by the *B.P.* The *P.G.* requires 24.95 p c and the *U.S.P.* not less than 30 p c of available Chlorine, the latter Pharmacopœia triturating a definite weight of the substance with Water, making up to a standard volume and performing the assay on an aliquot portion

Volumetric Determination—The *P.G.* uses 0.5 gramme Calcaria Chlorata mixed with a solution of 1 gramme Potassium Iodide in 20 c c of Water, and

acidulated with 20 drops Hydrochloric Acid, and directs that it should require at least 85 c c Tenth-normal Volumetric Sodium Thiosulphate Solution to combine with the liberated Iodine, *P G*

Introduce into a stoppered weighing bottle between 3 and 4 grammes of Chlorinated Lime and weigh accurately, titrate this thoroughly with 50 c c of mixture to a graduated vessel, together with the rinsings, and to make 1000 c c. After thoroughly shaking add to 100 c c of the mixture 1 gramme of Potassium Iodide, 5 c c Diluted Hydrochloric Acid and sufficient Tenth-Normal Volumetric Sodium Thiosulphate Solution for complete decolouration. Multiply the number of c c of Tenth normal Volumetric Solution consumed by 0.8518, and divide the product by one tenth of the weight of the Chlorinated Lime taken, the quotient represents the percentage of available Chlorine present, *U S P*

Preparations

LIQUOR CALCIS CHLORINATÆ. SOLUTION OF CHLORINATED LIME

Chlorinated Lime, 1, Distilled Water, 10 (1 in 10)

The Chlorinated Lime should be thoroughly mixed with the Water, and set aside in a stoppered bottle or three hours, shaking it at intervals, and finally strained.

yield, when fresh, about p c of available Chlorine, but on keeping. It should be preserved in well-stoppered amber-coloured bottles and kept in a cool and dark place.

Medicinal Properties.—A powerful antiseptic and bleaching agent. Diluted 1 to 12 or 16 of Distilled Water, it is used as an antiseptic lotion for unhealthy ulcers, purulent ophthalmia, iteal cutaneous diseases, as an injection in foul nasal, aural and as a gargle in septic tonsillitis and diphtheria.

Antidotes.—Emetics, White of Egg, Milk, Flour, not Acids.

Foreign Pharmacopœias.—Official in Belg, 3 in 100, Russ (Calcium Hypochlorosum Solutum), 2.5 p c of Chlorine, Span, about 1 in 48. Not in the others.

Tests.—The tests for Chlorinated Lime Solution are that it has a specific gravity of about 1.055 and shall yield not less than 2 p c of available Chlorine as determined by titration of the Iodine liberated, when 1 gramme of the liquor is mixed with 0.5 gramme of Potassium Iodide dissolved in Water and 1 c c of Diluted Hydrochloric Acid added, not less than 5.6 c c of Deci-normal Volumetric Sodium Thiosulphate should be required, corresponding to 1.97 p c of available Chlorine.

LIQUOR SODÆ CHLORINATÆ.—SOLUTION OF CHLORINATED SODA

An almost colourless liquid possessing an alkaline reaction, an astringent taste and a faint chlorinaceous odour. It is prepared by well rubbing 4 of Chlorinated Lime with 30 of Distilled Water and mixing this with 6 of Sodium Carbonate previously dissolved in 10 of Distilled Water, and filtering.

The method of preparation adopted in the *BP* 1885 recommended the solution of the Chlorinated Lime to be filtered, the solutions to be

well mixed and again filtered. The *Companion*, 1894 edition, pointed out that this was unadvisable, and the method was altered in accordance with the *Companion* recommendation in the 1898 edition of the *B P*.

It should be kept in well-stoppered amber-tinted glass bottles and in a cool and dark place. It has the reputation of being an unstable solution, but this is an error. It undergoes but slight change, even when kept under ordinary conditions during several months, or even after keeping for a week in an open white glass bottle.

The Labarraque Solution was prepared by mixing together the unfiltered solutions of one part of Chlorinated Lime with 2 parts of Soda crystals.

The proportions used in U S and other Pharmacopœias will be found below.

Medicinal Properties—Antiseptic. Used internally in typhoid fever and in dysentery. Invaluable as a gargle in throat affections attended with fœtor, as in scarlet fever, diphtheria and septic tonsillitis, 1 fl oz in 12 to 16 fl oz of Water. Diluted with Water or Glycerin it forms an excellent application to sore nipples. It is also a powerful disinfecting agent, and is employed as a wash for foul ulcers.

For information on the treatment of typhoid and diphtheria by Chlorine see under 'Chlorine Liquor'.

A paper by Klein on the disinfecting action of solutions of Sodium Hypochlorite—*L* '96, ii 1509.

Dose—10 to 20 minims = 0.6 to 1.2 cc

Foreign Pharmacopœias—Official in Fr (Chlorure de Soude dissous), Chlorinated Lime 1, Sodium Carbonate 2, Water 45, Mex (Hipoclorito de Sodio liquido), Sodium Chloride 8, Manganese Dioxide 8, Sulphuric Acid 8, Sodium Carbonate 5, Distilled Water 20, Port (Solutio de Soda Chlorada), Calcium Hypochlorite 1, Sodium Carbonate 2, Water 40, Span (Solucion de Hipoclorito Sódico), Calcium Hypochlorite 1, Sodium Carbonate 2, Water 43, Swiss (Natrium Hypochlorosum Solutum), Calcium Hypochlorite 4, Sodium Carbonate 5, Water 120, U S, Chlorinated Lime 90, Monohydrated Sodium Carbonate 65, Water to weigh 1000. Not in the others.

Tests—Chlorinated Soda Solution has a specific gravity of about 1.054, it decolorises Indigo Sulphate Solution, and yields when acidified with Hydrochloric Acid a yellowish-green gas possessing a strong chlorinaceous odour and which first reddens and then bleaches moistened blue Litmus paper. It is officially required to indicate at least 2.51 pc of available Chlorine as determined by the titration of the Iodine liberated when 3.5 grammes of the solution are added to a solution of 1 gramme of Potassium Iodide in 100 cc of Water and the mixture acidulated with 3 cc of Hydrochloric Acid, at least 25.0 cc of Volumetric Sodium Thiosulphate Solution should be required. The *USP* requires it to contain at least 2.4 pc by weight of available Chlorine, as volumetrically determined by mixing a weighed quantity of 7 grammes of the solution with 50 cc of Water and 2 grammes of Potassium Iodide, adding 10 cc of Hydrochloric Acid and titrating the liberated Iodine with Tenth-

normal Volumetric Sodium Thiosulphate Solution, of which not less than 48 c.c. should be required, 1 c.c. of Tenth-normal Volumetric Sodium Thiosulphate Solution is equivalent to 0.05 p.c. of available Chlorine. The preparation is not official in the *P. G.*

The more generally occurring impurities are Calcium and Carbonates. When acidified with Acetic Acid and warmed till Chlorine vapours cease to be evolved, it should not yield a pronounced turbidity on the addition of Ammonium Oxalate Solution, in the absence of more than a trace of Calcium, the gas evolved when the liquor is acidified with diluted Hydrochloric Acid should not cause a turbidity when passed into Lime Water, in the absence of Carbonates.

Not Official.

LIQUOR POTASSÆ CHLORINATÆ (Eau de Javelle) — Prepared by the interaction of Bleaching Powder and Potassium Carbonate. Contains about 3 p.c. available Chlorine.

CALX SULPHURATA.

SULPHURATED LIME.

Syn. — CALCIUM SULPHIDUM.

A white, or greenish-white amorphous powder possessing a characteristic odour of Hydrogen Sulphide. Should contain not much less than half its weight of Calcium Sulphide CaS , eq. 71.53, with Calcium Sulphate and Carbon.

It may be prepared by the reduction of native Calcium Sulphate by Carbon.

It should be kept in amber-tinted glass stoppered bottles and in a cool and dry place, as it is gradually decomposed by exposure to moist air.

Medicinal Properties — Antisuppurative, internally for boils, pustules and abscesses. In the form of Pigmentum or Lotio Calci Sulphurati for the cure of scabies, also used as a depilatory.

Daily doses of 1 grain as a prophylactic of influenza — *B. M. J.* '95, 1975.

Dose. — $\frac{1}{4}$ to 1 grain = 0.016 to 0.065 gramme.

Prescribing Notes — Best prepared in pill, made with Glucose. If the total weight of each pill be 500 grains, $\frac{1}{4}$ grain may be made up to this weight with Milk Sugar. The pills are coated with Sandarach solution, and usually sent out in bottles.

Foreign Pharmacopœias — Official in Jap., Mex., Port. and U.S. Not in the others.

Tests. — The following tests for Calcium Sulphide are that when acidulated with Acetic Acid it evolves a gas having a powerful odour of Hydrogen Sulphide, leaving a residue of Calcium Sulphate and Carbon, and when filtered the filtrate yields with Ammonium Oxalate solution a white precipitate insoluble in Acetic, but soluble in Hydrochloric Acid. Both *B.P.* and *U.S.P.* employ the Copper Sulphate test as a means of determining the presence of a due proportion of Sulphide, the *B.P.* requiring that a solution of 1.4 grammes of

Copper Sulphate in 50 c c of Water shall, when acidified with Hydrochloric Acid and brought nearly to the boiling point, be completely precipitated by 0.8 gramme of Calx Sulphurata, indicating not much less than 50 p c of pure Calcium Sulphide, the *U S P* that a solution of 1.9 grammes of Cupric Sulphate, when treated as indicated below shall be completely precipitated by 1 gramme of the substance, indicating the presence of at least 55 p c of pure Calcium Sulphide. The *B P* employs Potassium Ferriocyanide Solution, the *U S P* excess of Ammonia Water to detect excess of Copper Calx Sulphurata is not official in the *P G*.

The Zinc process described under Barium Sulphide is also applicable to Calx Sulphurata.

Determination—The *U S P* gives a similar test, using 1 gramme of Sulphurated Lime with a cold solution of 1.9 grammes of Copper Sulphate in 50 c c of Water and the addition of 10 c c diluted Hydrochloric Acid in small portions. The mixture is directed to be digested on a water bath for 15 minutes and filtered. The addition of an excess of Ammonia Water to the filtrate should impart no colour to it.

Not Official

SOLUTIO CALCII OXYSULFURATI (Solutio Vleminghx) —1 of Calcium Oxide slaked with 1 of Water, and mixed with 2 of washed Sulphur of the foregoing mixture 2.5 is boiled with 20 of Water until it is so reduced as to yield 10 by weight when strained.—*Aust*

1 of Calcium Oxide is treated with 5 of Water, and 2 of washed Sulphur with 15 of Water, mix and boil for one minute, when cold filter and wash the residue with Water to yield 12.—*Swiss*

Calcium Sulfuratum Solutum—Calcium Oxide, 10, Sulphur, 25, Water, 100.—*Belg*

Various formulas have been given for Vleminghx's Solution the proportion of Calcium Oxide, Sulphur and finished product varies between 2, 4, 20, 3, 5, 20, 2.5, 5, 20, 4, 4, 20.

Lotio Calcii Sulphurati—Calcium Hydrate, 4, Sublimed Sulphur, 4, Water, 85. Boil together, evaporate and filter, to produce 20 of solution. To be diluted with an equal quantity of warm Water.—*Westminster*

Liquor Calcis Sulphuratæ—Quicklime, 2, Sublimed Sulphur, 5, Water, q s to produce 100.—*B P G*

CAMBOGIA.

GAMBOGE

FR., GOMMEGUTTE; GER, GUMMIGUTT, ITAL, GOMMAGOTTA, SPAN, GUTAGAMBA

A gum-resin obtained from *Garcinia Hanbun*, Hook f

It is imported from Siam, and consists of about 75 p c of Resin and 15 to 20 of Gum, the Resin being the active ingredient.

Indian Gamboge, obtained from *Garcinia Morella*, Desr, is official in the *Ind* and *Col Add* for India and the Eastern Colonies.

Solubility.—About three-fourths is soluble in Alcohol (90 p c), the solution is rendered an opaque yellow by Water, three-fourths is also soluble in Ether. A solution in Ammoniated Alcohol is not rendered turbid by the addition of Water.

Medicinal Properties.—A powerful hydragogue cathartic, in small doses, diuretic. It is employed in the treatment of dropsy, attended with obstinate constipation, and in cerebral congestion. As it is apt to occasion much sickness and griping, it is best given in small doses, repeated at short intervals, until it operates, but it should never be given to children or very old persons, or in inflamed conditions of the abdominal or pelvic organs.

Dose.— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Ph Ger maximum single dose, 0.3 gramme, maximum daily dose, 1.0 gramme

Prescribing Notes—*It may be given in pill or emulsion, or dissolved in an alkaline solution, the last method has been recommended in dropsy*

Official Preparation—*Pilula Cambogiæ Composita*

Foreign P^r —Official in Aust^r (*Gummiresina Gutti*), Belg (*Guttiæ*) or (*Gutti*), Ital, Mex (*Goma Guta*), Port (*Gomma Guta*), Span, Swed (*Gummi Resina Gutta*), Swiss, U.S. (*Cambogia*) Not in the others

Descriptive Notes—It usually presents the cylindrical shape of the bamboo joints in which it is collected, it may be solid or hollow in the centre. If of good quality it has a smooth, even fracture, free from grittiness, of a bright orange colour. Inferior qualities have a dull and sometimes gritty fracture. The same remarks apply to Indian Gamboge.

Tests.—The distinguishing tests for Gamboge are that when rubbed with Water it forms a yellow emulsion, that it should be completely dissolved by the successive use of Alcohol (90 p.c.) and of Water. It has been suggested (*PJ '02*, n 495) that 75 p.c. should be soluble in Alcohol (90 p.c.), which agrees with the statement previously made in the *Companion* that three-fourths should be soluble in Alcohol (90 p.c.) This standard has been adopted by the *USP* using Alcohol (94.9 p.c.), but *PG* makes no reference to an Alcohol solubility.

The more generally occurring adulterants are Starch, mineral matter, and vegetable debris. Starch may be detected by the addition of Iodine to the cooled aqueous decoction, mineral matter by the ash left on incineration and vegetable matter by the increased insolubility in Alcohol (90 p.c.) The *BP* and *USP* require that when incinerated it shall yield not more than 3 p.c. of ash, the *PG* not more than 1 p.c.

Gamboge of good commercial quality contains from 60 to 80 p.c. of Resin (Gambogic Acid), from 15 to 25 p.c. of Gum, about 4 p.c. of Wax, moisture, mineral matter and a trace of Starch.

The *BP* states that it is completely dissolved by the successive action of Alcohol (90 p.c.) and Water, and the *USP* that not more than 25 p.c. should be insoluble in Alcohol (94.9 p.c.)

Preparation.

PILULA CAMBOGIÆ COMPOSITA. COMPOUND PILL OF GAMBOGI

Gamboge, 1, Barbados Aloes, 1, Hard Soap, 2, all in powder,

Compound Powder of Cinnamon, 1, massed with Glucose Syrup
(about 1 in 6)

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

Foreign Pharmacopœias—Official in Fr (Pilule Anderson), Aloes, Gamboge, Oil of Anise, and Honey, Poit (Pilulas de Aloes e Gomma Guta), the same with Soap, Mex (Pildoras de Anderson), Aloes 8, Gamboge 8, Oil of Anise 0.4, Soap 4, Water *q s*, also (Pildoras de Boncio), Aloes 7, Gamboge 7, Ammoniacum 7, Soap 4, Water *q s*, U S (Pil Cathartice Comp), contains Gamboge about 1 in 12 (see Colocynth) Not in the others

CAMPHORA.

CAMPHOR

FR, CAMPHRE DU JAPON, GER, KAMPHER, ITAL, CANFORA, SPAN, ALCANTOR

Refined Camphor is a white or colourless translucent crystalline solid. It is obtained in the raw state from *Cinnamomum Camphora*, Nees and Ebern, in Formosa and Japan, it is resublimed in this country and elsewhere.

It may also be produced synthetically by the oxidation of Camphene. Camphene exists in many essential oils, but commercially it is obtained from Pinene Hydrochloride or Borneol Hydrochloride by treatment with Alcoholic Potassium Hydroxide.

On account of its volatility it should be kept in well closed vessels, and in a cool place.

Borneo Camphor (Borneol) is a solid substance obtained from *Dryobalanops aromatica*, Gartin, in Borneo and its neighbourhood. Borneol is used in the production of artificial Camphor, two compound esters are known, Borneol Salicylate (Salit) and Borneol Valerianate (Bornyval).

Solubility—1 in 700 of Water, 1 in $1\frac{1}{4}$ of Alcohol (90 p c), or by weight, 1 in 1, 4 in 1 of Chloroform, 12 in 7 of Ether, 1 in 4 of Olive Oil (slowly), 1 in $1\frac{1}{2}$ of Oil of Turpentine, 2 in 1 of Glacial Acetic Acid, insoluble in alkalis.

3 of Camphor rubbed with 1 of Carbolic Acid crystals form a clear solution. 3 of Camphor and 3 of Chloral Hydrate rubbed together liquefy. Camphor also forms a liquid when mixed with many other substances, Menthol, Thymol, Naphthol, Salol, Butyl Chloral, and Salicylic Acid.

Medicinal Properties—A stimulant sedative, antispasmodic, expectorant, diaphoretic, and anaphrodisiac. A local anæsthetic. A feeble antiseptic.

Stimulant in the prostration of febrile diseases, sedative in mania, delirium tremens and chorea, also useful in dysmenorrhœa, spasmodic asthma and chronic bronchitis, in hysteria, nymphomania and spermatorrhœa. Spirit of Camphor mixed with warm Water to bathe the nostrils is highly useful in hay fever, and relieves irritation of the nostrils in common cold, also used as an inhalation. The **Compound Tincture** is given with Tincture of Squill to allay spasmodic cough in bronchitis and phthisis. In large doses Camphor tends to cause cardiac depression, convulsions, and possibly collapse.

Externally, it is used as a counter-irritant to relieve pain in chronic rheumatism, neuralgia, and as an application to chilblains, also in chronic eczema and other painful skin diseases. The combination with Thymol, Phenol, or Chloral forms a good local anodyne for neuralgia.

10 grammes of 10 p.c. solution of Camphor in Olive Oil hypodermically injected for collapse.—*B.M.J.E.* '95, ii 68, *P.J.* '95, ii 380.

Dose.—2 to 5 grains = 0.13 to 0.32 gramme

Prescribing Notes.—*An excellent pill can be made by mixing Camphor, 36 grains, Curd Soap, 4 grains, 'Diluted Glucose,' 10 grains, and dividing into 12 or more pills as required. Its unpleasant taste is covered well by Milk, which is a good solvent for Camphor. The Spirit is given on Sugar, also in Milk. Camphor can be powdered quite readily with the addition of a small quantity of Alcohol (90 p.c.)*

Symptoms of poisoning by Camphor: convulsions, lividity, stupor, arrest of urinary secretion.

Official Preparations.—Aqua Camphoræ, Linimentum Camphoræ, Linimentum Camphoræ Ammoniatum, Spiritus Camphoræ and Tinctura Camphoræ Composita. Contained in Linimentum Aconiti, Linimentum Belladonnæ, Linimentum Opii, Linimentum Saponis, Linimentum Sinapis, Linimentum Terebinthinæ and Unguentum Hydrargyri Compositum. Of **Linimentum Camphoræ**: Linimentum Chloroformi, Linimentum Hydrargyri, Linimentum Terebinthinæ Aceticum.

Not Official.—Aqua Camphoræ Conc., Camphor Ball, Camphora cum Creta, Camphorated Quinine, Ceratum Camphoræ, Essentia Camphoræ, Spiritus Camphoræ, or, Æther Spirituosus Camphoratus, Spiritus Camphoræ Compositus, Vinum Camphoratum, Essential Oil of Camphor, Eau de Cologne, Salicylate, Oxycamphor, Phenol Camphor, Thymol Camphor, Resorcin Camphor, and Camphoric Acid.

Antidotes.—Stomach-tube or emetics, stimulants freely, and warmth to the extremities.

Foreign Pharmacopœias.—Official in all.

Descriptive Notes.—It occurs in commerce in the form of 'Bells,' in rectangular pieces (Camphor squares) or in a pulverulent condition (Flowers of Camphor). It possesses a characteristic, powerful and penetrating odour and a pungent aromatic taste, subsequently producing a feeling of coldness in the mouth. It is described by the *U.S.P.* as the dextrogyrate modification of the saturated ketone $C_{15}H_{11}CO$. Squares containing Stearic Acid have been met with on the Continent. Genuine Camphor of Japanese manufacture has often an odour of Saffrol, which is less noticeable in Chinese or Formosa Camphor. Artificial Camphor has usually a faintly terebinthinate odour, and is liable to contain traces of Chlorine. Recently under the name of Pearl Camphor an artificial Camphor made in Germany and having a very pure odour has been offered in commerce in the form of small cylindrical pieces about $\frac{1}{8}$ in (4 mm) in diameter.

Tests.—The following tests for Camphor are its distinctive odour and physical appearance, the specific gravity (which should be 0.995), the melting point, which should be $175^{\circ}C.$ ($347^{\circ}F.$), and the boiling point, which should be $205^{\circ}C.$ ($401^{\circ}F.$). The *B.P.* only gives the specific gravity, which it states should be about 0.995, the

USP gives 0.990 at 25° C (77° F), the *USP* and *PG* give the melting point as 175° C (347° F), the *USP* gives the boiling point as 204° C (399.2° F), the *PG* gives neither specific gravity nor boiling point

The more generally occurring impurity is mineral matter, which is detected by the residue left on ignition. Artificial or synthetic Camphor is sometimes used, and the *USP* includes a test for Chlorinated compounds, which are indicative of the synthetic article, the test which is peculiar to the *USP* is described below in small type under the heading of Silver Nitrate. The solution in Alcohol (90 p.c.) should be neutral to Litmus, should Stearic Acid be present the solution would be acid, and the quantity could be ascertained by determining the Acid value.

Silver Nitrate—The *USP* gives the following test to detect chlorinated products. If a small piece of Camphor be dropped into a small porcelain dish, and a clean beaker moistened on the inner surface with Distilled Water be inverted over the smaller dish immediately after igniting the Camphor, a part of the products of combustion will be absorbed by the Water, if the beaker be rinsed with a little Distilled Water and the liquid filtered, the filtrate should yield no turbidity upon the addition of a few drops of Silver Nitrate T.S.

Preparations

AQUA CAMPHORÆ CAMPHOR WATER

Dissolve 70 grains of Camphor in Alcohol (90 p.c.) *q.s.* to form $\frac{1}{2}$ fl. oz., add this gradually to 160 oz. of Distilled Water, with agitation to form a solution (1 in 1000)

The solution of the Camphor in Alcohol saves time and ensures a more uniform product. The alcoholic solution of Camphor may be kept ready for use.

Dose—1 to 2 oz. = 28.4 to 56.8 c.c. = $\frac{1}{18}$ to $\frac{1}{8}$ grain of Camphor

Foreign Pharmacopœias—Official in Belg., Camphor 2, Alcohol 4, Water 994, Dan. (*Mistura Camphorata*), contains Camphor, Alcohol, Mucilage of Acacia, Syr. Cerasi and Water, Norw. (*Emulsio Camphoræ*), Camphor, Mucilage of Acacia, and Water, Port., Camphor 1, Water 100, Span. (*Emulsio Alcanforado*), Camphor, Sweet Almonds, Powdered Sugar, and Water, Swed. (*Emulsio Camphoræ*), Camphor, Gum Acacia, and Almond Emulsion, U.S., dissolve 8 of Camphor in 8 c.c. of Alcohol (95 p.c.) and triturate the solution with 15 of Purified Talc, and, after allowing the greater part of the Alcohol to evaporate spontaneously, continue the trituration with Distilled Water gradually added to make 1000 c.c., then pour the mixture upon a well wetted filter, and pass the filtrate through the filter repeatedly until the Camphor Water is perfectly clear. Not in the others.

LINIMENTUM CAMPHORÆ. LINIMENT OF CAMPHOR *B.P. Syn* —CAMPHORATED OIL

A yellow oily fluid, possessing a strong characteristic odour of Camphor, prepared by dissolving 1 of Camphor in flowers, in 4 of Olive Oil (about 1 in 5)

Solution will be more readily effected if the Camphor is sifted before using, and if the oil is warmed to about 38° C (100.4° F), agitating in a bottle or covered vessel to prevent vaporisation of the Camphor.

Foreign Pharmacopœias—*Off. and Disp., New York*, 1894, 1914, with Olive Oil, Belg., 1 and 9, *off. and Disp., Dutch*, 1914, 1916, 1918, 1920, 1922, 1924, 1926, 1928, 1930, 1932, 1934, 1936, 1938, 1940, 1942, 1944, 1946, 1948, 1950, 1952, 1954, 1956, 1958, 1960, 1962, 1964, 1966, 1968, 1970, 1972, 1974, 1976, 1978, 1980, 1982, 1984, 1986, 1988, 1990, 1992, 1994, 1996, 1998, 2000, 2002, 2004, 2006, 2008, 2010, 2012, 2014, 2016, 2018, 2020, 2022, 2024, 2026, 2028, 2030, 2032, 2034, 2036, 2038, 2040, 2042, 2044, 2046, 2048, 2050, 2052, 2054, 2056, 2058, 2060, 2062, 2064, 2066, 2068, 2070, 2072, 2074, 2076, 2078, 2080, 2082, 2084, 2086, 2088, 2090, 2092, 2094, 2096, 2098, 2100, 2102, 2104, 2106, 2108, 2110, 2112, 2114, 2116, 2118, 2120, 2122, 2124, 2126, 2128, 2130, 2132, 2134, 2136, 2138, 2140, 2142, 2144, 2146, 2148, 2150, 2152, 2154, 2156, 2158, 2160, 2162, 2164, 2166, 2168, 2170, 2172, 2174, 2176, 2178, 2180, 2182, 2184, 2186, 2188, 2190, 2192, 2194, 2196, 2198, 2200, 2202, 2204, 2206, 2208, 2210, 2212, 2214, 2216, 2218, 2220, 2222, 2224, 2226, 2228, 2230, 2232, 2234, 2236, 2238, 2240, 2242, 2244, 2246, 2248, 2250, 2252, 2254, 2256, 2258, 2260, 2262, 2264, 2266, 2268, 2270, 2272, 2274, 2276, 2278, 2280, 2282, 2284, 2286, 2288, 2290, 2292, 2294, 2296, 2298, 2300, 2302, 2304, 2306, 2308, 2310, 2312, 2314, 2316, 2318, 2320, 2322, 2324, 2326, 2328, 2330, 2332, 2334, 2336, 2338, 2340, 2342, 2344, 2346, 2348, 2350, 2352, 2354, 2356, 2358, 2360, 2362, 2364, 2366, 2368, 2370, 2372, 2374, 2376, 2378, 2380, 2382, 2384, 2386, 2388, 2390, 2392, 2394, 2396, 2398, 2400, 2402, 2404, 2406, 2408, 2410, 2412, 2414, 2416, 2418, 2420, 2422, 2424, 2426, 2428, 2430, 2432, 2434, 2436, 2438, 2440, 2442, 2444, 2446, 2448, 2450, 2452, 2454, 2456, 2458, 2460, 2462, 2464, 2466, 2468, 2470, 2472, 2474, 2476, 2478, 2480, 2482, 2484, 2486, 2488, 2490, 2492, 2494, 2496, 2498, 2500, 2502, 2504, 2506, 2508, 2510, 2512, 2514, 2516, 2518, 2520, 2522, 2524, 2526, 2528, 2530, 2532, 2534, 2536, 2538, 2540, 2542, 2544, 2546, 2548, 2550, 2552, 2554, 2556, 2558, 2560, 2562, 2564, 2566, 2568, 2570, 2572, 2574, 2576, 2578, 2580, 2582, 2584, 2586, 2588, 2590, 2592, 2594, 2596, 2598, 2600, 2602, 2604, 2606, 2608, 2610, 2612, 2614, 2616, 2618, 2620, 2622, 2624, 2626, 2628, 2630, 2632, 2634, 2636, 2638, 2640, 2642, 2644, 2646, 2648, 2650, 2652, 2654, 2656, 2658, 2660, 2662, 2664, 2666, 2668, 2670, 2672, 2674, 2676, 2678, 2680, 2682, 2684, 2686, 2688, 2690, 2692, 2694, 2696, 2698, 2700, 2702, 2704, 2706, 2708, 2710, 2712, 2714, 2716, 2718, 2720, 2722, 2724, 2726, 2728, 2730, 2732, 2734, 2736, 2738, 2740, 2742, 2744, 2746, 2748, 2750, 2752, 2754, 2756, 2758, 2760, 2762, 2764, 2766, 2768, 2770, 2772, 2774, 2776, 2778, 2780, 2782, 2784, 2786, 2788, 2790, 2792, 2794, 2796, 2798, 2800, 2802, 2804, 2806, 2808, 2810, 2812, 2814, 2816, 2818, 2820, 2822, 2824, 2826, 2828, 2830, 2832, 2834, 2836, 2838, 2840, 2842, 2844, 2846, 2848, 2850, 2852, 2854, 2856, 2858, 2860, 2862, 2864, 2866, 2868, 2870, 2872, 2874, 2876, 2878, 2880, 2882, 2884, 2886, 2888, 2890, 2892, 2894, 2896, 2898, 2900, 2902, 2904, 2906, 2908, 2910, 2912, 2914, 2916, 2918, 2920, 2922, 2924, 2926, 2928, 2930, 2932, 2934, 2936, 2938, 2940, 2942, 2944, 2946, 2948, 2950, 2952, 2954, 2956, 2958, 2960, 2962, 2964, 2966, 2968, 2970, 2972, 2974, 2976, 2978, 2980, 2982, 2984, 2986, 2988, 2990, 2992, 2994, 2996, 2998, 3000, 3002, 3004, 3006, 3008, 3010, 3012, 3014, 3016, 3018, 3020, 3022, 3024, 3026, 3028, 3030, 3032, 3034, 3036, 3038, 3040, 3042, 3044, 3046, 3048, 3050, 3052, 3054, 3056, 3058, 3060, 3062, 3064, 3066, 3068, 3070, 3072, 3074, 3076, 3078, 3080, 3082, 3084, 3086, 3088, 3090, 3092, 3094, 3096, 3098, 3100, 3102, 3104, 3106, 3108, 3110, 3112, 3114, 3116, 3118, 3120, 3122, 3124, 3126, 3128, 3130, 3132, 3134, 3136, 3138, 3140, 3142, 3144, 3146, 3148, 3150, 3152, 3154, 3156, 3158, 3160, 3162, 3164, 3166, 3168, 3170, 3172, 3174, 3176, 3178, 3180, 3182, 3184, 3186, 3188, 3190, 3192, 3194, 3196, 3198, 3200, 3202, 3204, 3206, 3208, 3210, 3212, 3214, 3216, 3218, 3220, 3222, 3224, 3226, 3228, 3230, 3232, 3234, 3236, 3238, 3240, 3242, 3244, 3246, 3248, 3250, 3252, 3254, 3256, 3

Tests.—Camphorated Oil has a specific gravity of 0.920 to 0.926, and should theoretically contain 21.45 p.c. w/w of Camphor, which may be determined by heating a definite weight of the sample in a flat-bottomed dish on a water-bath until it ceases to lose weight. Olive Oil is stated (*Analyst* xxiii 281) to suffer a gain in weight of 0.15 p.c. when heated for two hours at 120° C (248° F), and hence this figure should be added to the loss in weight when testing the samples under those conditions in order to obtain the true amount of Camphor present. But it has been subsequently shown (*C.D.* '01, 168) that it is best to make no such correction but to take the percentage of loss in weight as equal to the quantity of Camphor present.

Samples of the Liniment are frequently deficient in Camphor, and the loss of Camphor by volatilisation has been pleaded in justification of such deficiency, but it is generally conceded that if properly prepared it will not, when kept under ordinary conditions, lose any appreciable amount. The *USP* uses Cotton Seed Oil in the preparation of the Liniment, and Sesame Oil is employed by some of the Foreign Pharmacopoeias. Mineral Oil or a mixture of a mineral and vegetable oil has been used as an adulterant, but their presence is readily detected by determining the percentage of Potassium Hydroxide required to saponify the residue remaining after the volatilisation of the Camphor. The refractive index of the oil is apparently almost unaffected by the presence of dissolved Camphor (*Analyst* xxv, 202), and the refractometer may be employed for the identification of the oil used. It is shown (*CD '01, ii 390*) that the substitution of other oils for Olive Oil may, with the exception of Arachis Oil, be readily detected. The determination of the optical activity has also been suggested (*Analyst* xxv, 202) as a means of ascertaining the percentage of Camphor present in a sample. The rotation is increased by nearly 1° for each percentage of Camphor present, and the observed rotation of a sample in a 200 mm tube gives at once, without calculation, the percentage of Camphor with sufficient accuracy for most purposes. It has been pointed out (*CD '01, i 167*) that they show a slight over-estimation, the error varying with the amount of Camphor present, when 25 p.c of Camphor is present the division will be 0.998, while with 1 p.c it will be 0.987.

LINIMENTUM CAMPHORÆ AMMONIATUM. AMMONIATED
LINIMENT OF CAMPHOR *B.P. Syn* — COMPOUND LINIMENT OF
CAMPHOR *N.O. Syn* — LINIMENTUM AMMONIATUM CAMPHORATUM

Camphor, 5, Oil of Lavender, $\frac{1}{2}$; Strong Solution of Ammonia, 10, Alcohol (90 p c), *q s* to make 40. (1 in 8)

Dissolve the Camphor and the Oil in a portion of the Alcohol (about 24) and add the Ammonia gradually with agitation. The Camphor which is at first thrown out will readily redissolve. Add Alcohol *q s* to make 40.

Rubefacient and counter-irritant. Most useful in the douloureux and chronic rheumatism. Painful neuralgia is relieved by applying lint previously soaked in the liniment, covering with a dry napkin until redness is produced, then lightly rubbing the part with Laudanum or Liquor Meconicus.

• Foreign Pharmacopœias—Official in Belg (*Linimentum Ammoniacale Camphoratum*), Liquid Ammonia 1, Camphorated Oil 9, also (*Camphoræ Linimentum Compositum*), Spirit of Soap 700, Spirit of Camphor 250, Liquid Ammonia 30, Oil of Rosemary 15, Oil of Thyme 5, Dan, Solution of Ammonia 5, Camphor 1, Rape Oil 14, Fr, Solution of Ammonia 1, Camphorated Oil 9, Ger, Solution of Ammonia 1, Camphorated Oil 3, Poppy Oil 1, Ital, Solution of Ammonia 1, Camphorated Oil 4, Mex, Solution of Ammonia 1, Camphorated Oil 9, Norw, Solution of Ammonia 2, Camphorated Oil 1, Rape Oil 2, Port, Liquid Ammonia 1, Camphorated Oil 4, Russ, Solution of Ammonia 1, Camphorated Oil 3, Sesame Oil 1, Span, Solution of Ammonia 1, Camphorated Oil 9, Swed and Swiss, Solution of Ammonia 1, Camphorated Oil 3. All by weight. Not in the others.

Linimentum Ammoniacum Camphora—Solution of Ammonia, 25, Liniment of Camphor, 25, Olive Oil, *q s* to produce 100—*B P C Supplement*

SPIRITUS CAMPHORÆ SPIRIT OF CAMPHOR *N O Syn* — TINCTURA CAMPHORÆ

Camphor, 1, Alcohol (90 p c), *q s* to make 10 (1 in 10)

Dose—5 to 20 minims = 0.3 to 1.2 c c

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Dutch (*Solutio Camphoræ Spirituosa*), Fl (*Teinture de Camphré Conc*), Ger, Ital, Jap, Norw, Port, Swed, Swiss and US, 1 in 10, Hung, about 1 in 7, Russ, 1 in 13, Mex (*Alcohol Alcanforada*), 1 and 19, Span, 1 and 24, Fr has also a *Teinture de Camphre faible*, Camphor 1, Dilute Alcohol 39. All by weight except US.

Tests—Spirit of Camphor has a specific gravity of 0.843 to 0.847. It should contain 10 p c w/v of Camphor, which may be determined by its optical rotation.

A paper on the determination of Camphor in Spirit of Camphor appears in *CD* '99, 154.

A process requiring less manipulative skill, and which is claimed to give very exact results, appears in *YBP* '01, 47. A weighed quantity of 10 grammes of the spirit is introduced into a 50 c c burette, graduated in $\frac{1}{10}$ c c and shaken with 30 to 35 c c of a saturated solution of Sodium Chloride. 1 c c of Petroleum Ether is dropped on the Camphor layer when it has collected as much as possible on the surface, and solution of the Camphor effected by careful agitation. After several minutes the volume of the Benzin solution can be read off. After subtracting the volume of added Benzin each further 1.02 c c will correspond to 1 gramme of Camphor (sp gr 0.98).

TINCTURA CAMPHORÆ COMPOSITA. COMPOUND TINCTURE OF CAMPHOR *B P Syn*—PAREGORIC PAREGORIC ELIXIR

Tincture of Opium, 585 minims, Benzoic Acid, 40 grains,

Camphor, 30 grains, Oil of Anise, $\frac{1}{2}$ fl dm, Alcohol (60 pc), *qs* to make 20 fl oz

Dissolve the Benzoic Acid, Camphor, and Oil of Anise in a portion of the Alcohol (about 17 or 18), add the Tincture and finally Alcohol *qs* to make 20 fl oz

B P 1898 replaced the 40 grains of Opium by the quantity of Tincture of Opium, 585 minims, as previously suggested in the *Compendium*

The Pimpinella Oil is preferable as being more soluble in Alcohol (60 pc)

Dose—30 to 60 minims = 1.8 to 3.6 cc = $\frac{1}{8}$ to $\frac{1}{4}$ grain Opium

30 minims contain about $\frac{1}{80}$ gram of anhydrous Morphine (about $\frac{1}{60}$ grain of

pc w/v of anhydrous Morphine
Opium Benzoica, the equivalent in the Foreign Pharmacopœias of our Compound Tincture of Camphor shall contain 0.05 pc w/v of Morphine Tinctura Camphora *Co B P* contains 0.05 pc (more correctly 0.046 pc) w/v of anhydrous Morphine

Official Tincture of Opium contains 0.75 pc of anhydrous Morphine

Foreign Pharmacopœias—Official in Belg (*Opium Tinctura cum Acido Benzoico*), Tincture of Opium 50, Benzoic Acid 5, Camphor 3, Anethol 2, Alcohol (70 pc), 940, Dan (*Tinctura Thebaïca Benzoica*), Tincture of Opium 50, Benzoic Acid 5, Camphor 3, Oil of Anise 2, Diluted Alcohol 940, Dutch (*Tinctura Opium Benzoica*), Tincture of Opium 10, Benzoic Acid 4, Camphor 2, Oil of Anise 1, Diluted Alcohol 188, Fr (*Elixir Paregorique*), Opium 5, Benzoic Acid 5, Camphor 2, Oil of Anise 5, Alcohol (60 pc), 985, Ger and Russ (*Tinctura Opium Benzoica*), Opium 1, Benzoic Acid 4, Camphor 2, Oil of Anise 1, Diluted Alcohol 192, Jap, Opium 1, Benzoic Acid 4, Purified Camphor 2, Fennel Oil 1, Diluted Alcohol 192, Norw (*Tinctura Opium Benzoica*) Tincture of Opium 50, Benzoic Acid 5, Camphor 3, Oil of Anise 2, Diluted Alcohol 940, Mex (*Tinctura de Opio Alecanforado*), Extract of Opium 3, Benzoic Acid 3, Camphor 2, Oil of Anise 3, Alcohol (60 pc) to 600, Port (*Tinctura de Opio Composita*), Opium 1, Benzoic Acid 1, Camphor 1, Oil of Anise 1, Alcohol (65 pc) 196, Swiss, Tincture of Opium 10, Benzoic Acid 1, Camphor 1, Oil of Anise 1, Diluted Alcohol 187, U S (*Tinctura Opium Camphorata*), Opium 4, Benzoic Acid 4, Camphor 4, Oil of Anise 4, Glycerin 40, Diluted Alcohol to 1000. All by weight, except U S. Swed (*Tinctura Opium Benzoica*), Opium 5, Benzoic Acid 5, Camphor 3, Oil of Anise 2, Diluted Alcohol to 1000. Belg, Fr and Swiss adopt the *Brussels Conference* standard

Tests.—Compound Tincture of Camphor has a specific gravity of 0.915 to 0.920, contains 0.2 to 0.35 pc w/v of total solids, the percentage of Absolute Alcohol should be about 58.0 pc by volume, 100 cc should contain nearly 0.05 gramme of anhydrous Morphine, it should afford well-marked reactions for Benzoic and Meconic Acids, and when diluted with Water should assume an opalescent appearance

It is frequently found deficient in Alcohol, in the requisite proportion of Tincture of Opium, the Benzoic Acid may be either of inferior quality or omitted, and Oil of Anise may be absent. The specific gravity of the sample affords a fair indication of the percentage of Absolute Alcohol, and may be supplemented if necessary by an actual determination of the Alcohol by distillation. The Benzoic Acid may be determined by rendering 10 cc of the Tincture alkaline by the addition of Sodium Hydroxide Solution and removing the Alcohol by evaporation. The aqueous liquid may then be extracted with Ether, the ethereal solution separated, the aqueous

fluid faintly acidified with Hydrochloric Acid and again shaken with Ether, the ethereal solution removed, the Ether evaporated spontaneously and the Benzoic Acid weighed, or the ethereal solution may be freed from mineral acid by carefully washing with Water and titrated with Twentieth-normal Volumetric Potassium or Sodium Hydroxide Solution. Each c.c. of Twentieth normal Solution represents 0.00605 gramme of Benzoic Acid. The separated Benzoic Acid should possess the melting point, and should otherwise correspond to the tests given under Benzoic Acid. Meconic Acid may be detected by diluting a little of the Tincture with Alcohol (45 p.c.) until of a pale straw colour and adding a drop of Ferric Chloride Test-solution, when a deep red coloration will be produced. The quantity of Morphine present is too small to permit the employment of the *B.P.* method of determination. A process depending on the extraction of the Morphine from the evaporated Tincture by hot Amylic Alcohol is described (*Y.B.P.* '05, 461, *Analyst*, '05, 336), the extracted Morphine being subsequently recognised by Ferric Chloride Test solution and the Nitric Acid colour test. It has been pointed out by Allen (*Analyst*, '02, 352) that cough mixtures, in addition to containing Paregoric, frequently contain Ipecacuanha in some form or other, and the alkaloids of the latter are liable to be mistaken for Morphine if colour reactions alone are relied upon. These colour reactions are given under the heading of Ipecacuanha and Psychotrine. A process for the analysis of the Tincture is recorded, and the use of hot Amylic Alcohol for the extraction of the Morphine is suggested in the paper.

Not Official

AQUA CAMPHORÆ CONC—Spirit of Camphor, 3 fl. dim., Distilled Water, 40 fl. oz.—*Pharm. Form (MacEwan)*

CAMPBOR BALL—Camphor, 2, White Beeswax, 5, Spermaceti, 3, Oil of Almonds, 8, Tincture of Tolu, $\frac{1}{2}$, melt and pour into $\frac{1}{2}$ oz. moulds.

The proportion of the Wax to Spermaceti varies in different forms, but the total of the two combined is generally much the same.

CAMPFORA CUM CRETA (Camphorated Chalk)—Camphor, 1, Precipitated Chalk, 8, powder the Camphor by rubbing it with a few drops of Alcohol (90 p.c.), mix in the Chalk, and pass the whole through a sieve.

This is used as a dentifrice, it is also made in the proportion 1 to 7 (*Mart*) and 1 to 9 (*B.P.C.*)

CAMPFORATED CHLOROFORM See **CHLOROFORM**

CAMPFORATED QUININE—Camphor, in powder, 8 grains, Ammoniated Tincture of Quinine, *q.s.* to make 1 fl. oz.

A very useful combination for an ordinary cold in the head. Dose—1 to 2 fl. dim.

Camphorated Quinine Capsules—Quinine Sulphate, 60 grains, Ammonium Carbonate (powdered finely), 100 grains, Powdered Camphor, 50 grains, Soft Paraffin and Liquid Paraffin, *q.s.* to make a thin paste and fill 100 capsules. Each capsule represents about 30 minims of Camphorated Quinine.

CERATUM CAMPHORÆ—Camphor, 2, White Beeswax, 8, Benzoinated Lard, 4, Oil of Almonds, 8, melt together and stir till cold.

U.S.P. **CERATUM CAMPHORÆ** is much weaker in Camphor—Camphor Liniment, 10, White Wax, 35, White Petrolatum, 15, Benzoinated Lard, 40, it contains 1 of Camphor in 50.

Belg (Unguentum Camphoratum), Camphor 1, Simple Ointment 4;
Fr (Pommade Camphrée), Camphor 2, Benzoinated Lard 7, White Wax
1, Mex (Pomada Alcanfoiada), Camphor 1, White Wax 1, Lard 8
B P C Unguentum Camphoræ—Camphor, 1, Soft Paraffin, 9
B P C Unguentum Camphoræ Durum (*Syn* Camphor Ice)—Camphor,
10, Hard Paraffin, 22, Soft Paraffin, 68

PHENOL CAMPHOR See ACIDUM CARBOLICUM

ESSENTIA CAMPHORÆ—Camphor, 1, Alcohol (90 p c), 20 Given for
coryza, 5 minims every hour in Water or on Sugar

This is half the strength of the official *Spiritus Camphoræ*

SPIRITUS CAMPHORÆ FORTIOR (Rubini's Essence)—A saturated
solution, in Alcohol (90 p c), contains about 1 in 2½

This has been incorporated in the *B P C* as **Essentia Camphoræ**, with the
names *Spiritus Camphoræ Fortior*, and Rubini's Essence of Camphor as synonyms,
as follows—Camphor, 2, Alcohol (90 p c), *q s* to make 5

ÆTHER SPIRITUOSUS CAMPHORATUS—Camphor 15, Spirit of
Ether 85—*Dan*, *Norw* and *Swed*

SPIRITUS CAMPHORÆ COMPOSITUS (*B P C*)—Camphor, 0 25,
Benzoic Acid, 0 50, Oil of Anise, 0 25, Liquid Extract of Liquorice, 1, Alcohol
(60 p c), *q s* to produce 100

SYRUPUS CAMPHORÆ COMPOSITUS—Benzoic Acid, 3 drm,
Glacial Acetic Acid, 3 oz 5 drm 20 minims, Vinegar of Squill, 2 pints, Vinegar
of Ipecacuanha, 40 fl oz, Anise Oil, 2 drm, Camphor, 2 drm, Tincture of
Opium, 10 oz 5 drm 20 minims, Sugar, 28 lb, Burnt Sugar, *q s*, Water, to
make 4 gall Dose—1 drm (= 1 minim of Tincture of Opium)—*Bristol Royal
Infirmary*

This has been incorporated in the *B P C*

CAMPHOID (Martindale)—Camphor, 20, Absolute Alcohol (100 p c) 20,
Pyroxylin, 1 Employed as a solvent for Iodoform, the odour of which it is said
to disguise—*P J '92*, 1 831, *B M J '92*, 1 1086

VINUM CAMPHORATUM (*Ger*)—Camphor, 1, Alcohol (90 p c), 1,
Mucilage of Acacia, 3, White Wine, 45 *Swiss*, Camphor 2, Alcohol (90 p c) 3,
Acacia 2, White Wine 93 All by weight

ESSENTIAL OIL OF CAMPHOR—Camphor Oil as it is distilled from
Camphor Wood contains some quantity of solid Camphor which is separated by
filtration and sometimes by refrigeration It has been used as an
application in

Two kinds of Camphor Oil are known commercially, Japan and China
Camphor Oil

Japan Camphor Oil—A yellow or yellowish-brown liquid possessing a
strong odour of Saffrol It has a specific gravity of 1 010 to 1 020, and an
optical rotation in a 100 mm tube of + 5 5° It dissolves readily in Alcohol
(90 p c)

A considerable portion of the Saffrol is frequently removed from the oil,
which is then commonly known in the trade as 'White Oil of Camphor' The
abstraction of Saffrol is indicated by a decrease in the specific gravity

China Camphor Oil—A pale yellow or brownish yellow liquid, possessing
a strong camphoraceous odour It has a specific gravity of 0 950 to 0 960, and
an optical rotation in a 100 mm tube of about + 30° It dissolves readily in
Alcohol (90 p c)

EAU SEDATIVE (*Lotion Ammoniacale Camphrée*, *Fr*)—Spirit
of Camphor, 10, Tincture of Camphor 60, Sodium Chloride, 60, Distilled Water, 1000
Belg, Sodium Chloride 60, Liquid Ammonia 60, Spirit of Camphor 10,
Water 870, *Mex*, Sodium Chloride 60, Liquid Ammonia 60, Spirit of Camphor
10, Water 1000, *Swiss*, Sodium Chloride 60, Water 880, Spirit of Camphor 10,
Liquid Ammonia 100

CAMPHOR SALICYLATE (Camphossil) —A crystalline, unctuous, deliquescent mass. Condensation product of Camphor and Salicylic Acid. Introduced as an antiseptic and antipyretic.

Dose — $7\frac{1}{2}$ grains = 0.5 gramme

OXYCAMPHOR —A white crystalline powder, soluble about 1 in 50 Water. Has been found useful in cases of dyspnoea, especially of pulmonary origin. Best given in cachets or gelatin capsules. It is easily altered by exposure to air. — *P J '02*, 11 132

Dose —15 to 30 grains = 1 to 2 grammes

Under the name of **Oxaphor**, a 50 p.c. solution of the above in Alcohol (90 p.c.) has been introduced. **Camphoroxol** is stated to be a 1 p.c. alcoholic solution of Camphor containing 3 p.c. Hydrogen Dioxide.

Thymol-Camphor and **Resorein-Camphor** are oily fluids obtained by heating Camphor with equal parts of Thymol and Resorein respectively.

CAMPHORIC ACID —Colourless, crystalline leaflets, or a white, crystalline powder, with a faint camphoraceous odour.

It is a di-basic acid prepared by the oxidation of Camphor. It should be kept in well closed vessels.

Solubility —1 in 160 of cold Water, 1 in 8 of boiling Water, 1 in 1.5 of Alcohol (90 p.c.), readily in Ether.

Dose —15 to 30 grains = 1 to 2 grammes, conveniently given in cachets.

Is a valuable remedy in cases of urinary calculi and of vesical catarrh. A 1 p.c. solution has been recommended in acute and chronic affections of the respiratory passages. — *P J (3) xix* 507

In 4 p.c. alcoholic solutions as spray or linctus, in laryngeal phthisis.

In cystitis, 15 grains 3 times a day. — *Y B P '02* 167

One gramme 3 or 4 times a day, or 2 grammes in the evening, checks the night sweating in phthisis. — *L M R '88*, 276

Foreign Pharmacopœias —Official in Dutch, Ger., Jap., Swiss and U.S.

Tests —Camphoric Acid has a melting point of 186° to 187° C (366.8° to 368.6° F), and is dextro-rotatory, a 10 p.c. alcoholic solution showing a value $[\alpha]_D = +47.8^\circ$. The aqueous solution is acid in reaction towards blue Litmus paper. 1 gramme of the acid should require for neutralisation not less than 10.0 c.c. of Normal Volumetric Potassium or Sodium Hydroxide Solution, indicating not less than 99.81 p.c. of pure Camphoric Acid.

The more generally occurring impurities are Chlorides, Sulphates, Nitrates, and mineral matter. The saturated aqueous solution should not be rendered turbid by Silver Nitrate nor by Barium Chloride Solution. When a solution of Ferrous Sulphate is poured carefully on to the cold saturated aqueous solution of the acid mixed with an equal quantity of Sulphuric Acid, no dark coloration should be developed at the line of junction of the two fluids. It should leave no weighable residue after ignition.

Not Official

CAMPHORA MONOBROMATA

MONOBROMATED CAMPHOR

$C_{10}H_{16}BrO$, eq 229.38

FR., CAMPHRE MONOBROMÉ, GER., MONOBROMKAMPFER, ITAL., CANFORA MONOBROMATA, SPAN., ALCANFOR MONOBROMADO

Colourless, prismatic needles, or scales, with a camphoraceous odour and taste.

It is a substitution product of Camphor, the Hydrogen radicle of the latter being replaced by the halogen Bromine.

It should be kept in well closed bottles.

Sol. b. insoluble in Water; soluble 1 in 12 of Alcohol (90 p c);
101 in 2 of Ether, 1 in 8 of Olive Oil, sparingly in Glycerin

Medicinal Properties.—Hypnotic and sedative. Given in hysteria, epilepsy, chorea, spermatorrhœa, and delirium tremens, but its use requires caution. It has been stated to be an antidote to Styrchnine.

Dose—2 to 5 grains = 0.13 to 0.32 gramme

Prescribing Notes—It can be given in pills with 'Diluted Glucose,' or can be dissolved in Almond oil and emulsified with Mucilage and Water. It is also given with Extract of Belladonna. Larger doses than 5 grains are sometimes given in delirium tremens.

Foreign Pharmacopœias—Official in Dutch, Fr., Ital., Jap., Mex. (Alcanfor Monobromado), Port., Span. (Alcanfor Monobromado), Swiss and U.S. Not in the others.

Tests—Monobromated Camphor has a melting point of 76° C (168.8° F), and a boiling point of 274° C (525.2° F). It should be neutral in reaction towards Litmus paper. When boiled with Silver Nitrate Solution a precipitate of Silver Bromide is formed. It is soluble in cold concentrated Sulphuric Acid without alteration in colour and without decomposition, and is again precipitated when this solution is poured into Water. When fused with metallic Sodium in a dry test-tube and the residue dissolved in Water, the resulting solution acidified with Nitric Acid yields with Silver Nitrate Solution a copious, faintly yellowish precipitate almost insoluble in weak Ammonia Solution.

The more generally occurring impurities are Bromides, Hydrobromic Acid, and mineral residue. The presence of Bromides is readily detected by the formation of an opalescence or precipitate when treated with Silver Nitrate Solution, Hydrobromic Acid by the reaction towards blue Litmus paper. It should be completely volatilised by heat leaving no weighable residue, showing the absence of mineral matter.

CANNABIS INDICA.

INDIAN HEMP

FR., CHANVRE INDIEN, GER., INDISCHER HANF, ITAL., CANAPE INDIANA,
SPAN., CAÑAMO

The dried flowering or fruiting tops of the female plant of *Cannabis sativa*, Linn., from which, of course, the Resin has not been removed, grown in India.

O Shaughnessy introduced Indian Hemp into this country, and Peter Squire made the extract for him.

The official variety may consist, according to the official monograph, of either the flowering or fruiting tops, and is frequently of very inferior quality, since the fruiting tops yield less Resin.

Preparations of *Cannabis Indica* show marked variation in physiological activity, owing to age, and perhaps other causes.

It is suggested by Dr. Marshall, the drug and its preparations should be kept in hermetically-sealed packages if they are to be kept for any considerable time before use. The drug should be purchased once a year only, when the new harvest comes in. This is generally in April or May—*P. J.* '02, 1342, 362, 892, '02, 11 131, 263, 284, *C. D.* '02, 11 296, *Y. B. P.* '02, 52, 168, 401.

A description of the names which are applied to the different forms of the drug, e.g., Ganja, Guaza, Bhang, Chur, Churrus or Charas, Haschish and Majun, and useful hints regarding the collection and storage of the drug—*P. J.* '02, 11 131.

The important constituent is a Resin (Extract of Indian Hemp); the active principle is stated to be a red Oil, *Cannabinol*, which is liable to become

oxidised and inert The ethereal extract from Charas yielded four distinct chemical substances (1) about 1.5 p.c. of a Terpene, boiling at 160° to 180° C (320° to 356° F), (2) 2 p.c. of a Sesquiterpene, boiling at 258° to 259° C (496° to 498° F), (3) 0.15 p.c. of a Paraffin, $C_{26}H_{54}$, melting point, 63° to 64° C (145° to 147° F), (4) 33 p.c. of a toxic red oil, Cannabinol, $C_{21}H_{30}O_2$, boiling at 265° C (509° F), under a pressure of 20 mm.—*J C S Trans* '96, 539, '99, 20

A Cannabinol prepared according to Wood, Spivey and Eastenfield's process was found to be physiologically inactive, and it was therefore concluded, though apparently without sufficient reason, that it is not the active principle of Cannabis, and so far as present information extends, the active principle has not yet been isolated.—*C D '02*, n 296, *P J '02*, n 131, 171, *Y B P '02*, 399

Medicinal Properties—Sedative, anodyne, and antispasmodic. Has been used with success in migraine and delirium, neuralgia, pain of last stages of phthisis, and in acute mania, also in menorrhagia and dysmenorrhœa. It is combined with Belladonna in whooping-cough, and in infantile convulsions, hepatic and renal colic, it is given in tetanus and hydrophobia.

Prescribing Notes—Usually prescribed in the form of Extract or Tincture.

Dose of the Extract $\frac{1}{4}$ to 1 gram (with a sufficiency of Liquorice Powder to form a pill), but as it varies considerably in strength it is better to commence with the smaller dose, toxic symptoms have been produced with 1 gram. *Dose of the Tincture*, 5 to 15 minims, which can be taken on Sugar, or diffused in Water by the aid of 1 fl. dr. of Mucilage of Acacia to each fl. oz. of Water, the Mucilage should be diluted with twice its volume of Water before the addition of the Tincture.

Two interesting cases of toxic symptoms, caused in one case by taking the whole of the active ingredient of a mixture in the last dose, owing to omission of Mucilage for suspension. The other, a nervous patient, for whom the *B P* minimum dose was prescribed, and who took a dose from the bottle without measuring, and inadvertently took rather more than a double dose.—*L '03*, i 1042

Official Preparations—*Extractum Cannabis Indicæ*. Of the **Extract**, *Tinctura Cannabis Indicæ*. The **Tincture** is contained in *Tinctura Chloroformi et Morphinæ Composita*.

Not Official—*Cannabinæ Tannas*, *Cannabinon*, and *Fluidextractum Cannabis Indicæ*.

Antidotes—In case of over-dose, after employing stomach tube, or emetics, hot brandy and-water may be given, vegetable acids, such as lemon juice, vinegar, and the like. *Stychnine* should be injected and a blister applied to the nape of the neck.

Foreign Pharmacopœias—Official in Austr., Belg., Dutch, Hung., Ital., Jap., Norw. (*F. luctus Cannabis*), Port. (*Canhamo*), Russ., Mex. (*Marihuana*), Swed., Swiss and U.S. Not in Ger.

It does not produce constipation or loss of appetite, on the contrary, it restores the appetite which has been lost by chronic Opium and Chloral drinking.—*L '89*, i 625

In chorea and pertussis.—*L '02*, i 1159

Descriptive Notes—Indian hemp is imported from Bombay under the name of Guaza, in compressed or flattened masses, consisting of the flowering and fruiting tops of the plant, matted together with its resinous secretion. From Calcutta it is sometimes imported in a rolled or cylindrical form under the name of Ganja (*P J* (4) xv. 129, xi p 782) but this form, which is much more active, rarely remains in this country, being exported to the West Indies (*P J* (4) x p 522). The best quality of either kind is seldom exported from India, since the natives are aware that it rapidly loses its activity on

keeping and therefore retain the current crop for home use, exporting that of the previous year (*PJ* (4) xiv p 342) The form of the drug official in the *B.P* is the compressed form of Guaza The upper leaves which accompany the flower are alternate and 1-3 partite, whilst the lower leaves have 5-7 linear serrate leaflets and are opposite The bracts as well as the leaves have characteristic resin glands and one-celled curved hairs with a cystolith contained in the enlarged base of each The bract below the fruit is ovate-lanceolate

During the last few years compressed *Cannabis Indica* has been imported from Zanzibar, Delagoa Bay, and from the North of France, but according to Dr W E Dixon is not nearly so active as the Indian drug (*PJ* (4) xx p 550) This difference in strength is said to be due to the fact that the female plant yields less resin if male plants are present in the fields In India the male plants are always removed before the plant flowers by experts employed for that purpose

Tests.—Three typical samples of Ganja, collected at the proper season and imported direct from India, when examined in the author's laboratory yielded a total extract to Alcohol (90 p c) of 15.52, 15.77 and 16.85 p c, when evaporated to dryness and again dissolved in spirit the Alcohol-soluble matter amounted to 14.45, 14.75 and 16.7 p c, leaving 1.07, 1.02 and 0.68 p c of insoluble brown extract The ash of the 3 samples amounted respectively to 12.70, 12.30 and 11.90 p c

Preparations.

EXTRACTUM CANNABIS INDICÆ. EXTRACT OF INDIAN HEMP

Exhaust Indian Hemp, in coarse powder, with Alcohol (90 p c) by percolation, evaporate to a soft extract

Dose.— $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Hung, Ital., Port, Russ (Extracto de Marihuana), and Swiss and U S Not in the others U S has also *Fluidextractum Cannabis Indicæ* (1 in 1)

Commercial extracts have been shown (*PJ* '02, 1 234, 281, 301) to vary greatly in strength, and to contain a doubtful and variable amount of alcohol-soluble matter in Alcohol (90 p c) The use of an ethereal instead of the official extract, or, better still, one made with Absolute Alcohol, is suggested Spiritus Ætheris *B.P* is recommended as a menstruum for preparing the Tincture

TINCTURA CANNABIS INDICÆ. TINCTURE OF INDIAN HEMP.

Dissolve 1 of Extract of Indian Hemp in 18 of Alcohol (90 p c), filter if necessary, and add Alcohol (90 p c), *qs* to yield 20

(1 in 20)

22 minims contain 1 grain of Extract

Dose.—5 to 15 minims = 0.3 to 0.9 c c.

Foreign Pharmacopœias—Official in Jap and Port, 1 Extract in 20; the following are from Herb. Ind. Mex, 1 in 5, Russ and Swiss, 1 in 10, all by weight, U S, 1 in 10, and others.

Tests.—Tincture of Cannabis Indica has a specific gravity of 0.845 to 0.850, it contains about 4 p.c. w/v of total solids and about 87 p.c. w/v of Absolute Alcohol

Not Official

FLUIDEXTRACTUM CANNABIS INDICÆ (U.S.)—10 of Cannabis Indica, in No. 30 powder, exhausted by percolation with Alcohol (94.9 p.c.), reserving the first 9, evaporating the remainder (not above 50° C.) to a soft extract which dissolve in reserved portion, and add Alcohol (94.9 p.c.) q.s. to make 10

CANNABINÆ TANNAS—An amorphous, yellowish powder, sparingly soluble in Water, Alcohol, and Ether Soluble in acidulated Alcohol

Dose—4 to 8 grains = 0.26 to 0.52 gramme, mixed with Sugar and taken as a powder or in a cachet

Was introduced as a hypnotic, but its effects are very uncertain—*T. G.* '85, 329, 379

It is occasionally prescribed for menorrhagia

CANNABINON—A soft resinous substance, generally found as a 10 p.c. trituration with Milk Sugar, also introduced as a hypnotic, but the dose (1½ grains) was followed by excitement, collapse, and cramps—*T. G.* '85, 286, *L. M. R.* '86, 434, contra indicated in cardiac disease—*L.* '87, 1542

Dose—½ to 1 grain = 0.016 to 0.06 gramme

CANTHARIS.

CANTHARIDES

FR, CANTHARIDE, GER, SPANISCHE FLIEGEN, ITAL, CANTARIDE, SPAN, CANTARIDA

The dried Beetle, *Cantharis vesicatoria*, Latr

It is collected in Spain, France, Russia, Sicily, and Hungary

The powder should be dry and kept closely corked, for if at all damp it is apt to acquire a putrid odour A piece of Camphor kept in it prevents mites

Mylabris—The dried beetle, *Mylabris phalerata*, is official in the *Ind* and *Col. Add.* for India and the African and Eastern Colonies, other species of *Mylabris* may be used provided they yield an equal amount of Cantharidin

Medicinal Properties—Externally its effects are rubefacient and irritant, by continued application it is vesicant For the latter purpose the Emplastrum or Liquor Epispasticus is used, and is especially effective in inflammation of deep-seated parts, as in pleuritis, pericarditis, pneumonia, sciatica, neuralgia, and over the præcordial region in acute rheumatism, applied to rheumatic joints it removes pain and swelling, applied over the epigastrium it often checks obstinate vomiting and gastric pain It acts for a longer period, and is less irritating to the patient, than Ammoniacal or Acetic Acid embrocations It ought not to be applied to a paralysed limb Internally in small doses it is diuretic and aphrodisiac It is given in gleet, in impotence, and incontinence of urine due to paralysis, but it should be given cautiously, for it irritates the kidneys and sometimes produces strangury, and it should never be given to aged people or to children, or in cases of nephritis

Continuous counter-irritation of the spine by blisters in the neighbourhood of the cervical and lumbar enlargements the most successful treatment of rheumatoid arthritis—*L.* '07, 11895

The Tincture in 5-mm doses three times daily in Water arrests hæmorrhage from the kidney.—*B M J* '98, ii 1551

It is the basis of most of the applications used to increase the growth of hair

In chronic inflammation of the bladder it should *not* be used as a counter-irritant, on account of its irritating effects on the urinary organs, when absorbed. In such cases a solution of Silver Nitrate ($\frac{1}{2}$ drm to 1 fl oz of Water) is to be preferred

Thirty-two cases out of 56 of cystitis cured by teaspoonful doses of the following solution. Cantharidin, 1 milligramme, dissolved in 1 gramme of Alcohol, and diluted to 100 grammes with Water.—*B M J E* '95, ii 6

Ph Ger maximum single dose, 0.05 gramme, maximum daily dose, 0.15 gramme

Official Preparations—Acetum Cantharidis, Emplastrum Calefaciens, —
Cantharidis, Liquor Epispasticus, Tinctura Cantharidis, and
Cantharidis Colloidium Vesicans is prepared from Liquor Epispasticus

Not Official—Cantharidin, Potassium Cantharidate, Charta —
Emplastrum Vesicans, Linimentum Cinale, Liquor Cantharidis —
Unguentum Stimulans, and Boni's Blister.

Antidotes—Emetics or stomach-tube, followed by Barley Water, Gruel, white of Egg, inject Morphine for pain

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Descriptive Notes.—The dried beetle, *Cantharis vesicatoria*, Latr., has shining coppery green, or green coloured wing cases, and is of an oblong shape, 18 to 25 mm long (1.5 to 3 cm, and 6 to 8 mm broad, *P G*), and has an unpleasant odour. The insects or powder should be dried when received into stock (preferably in the presence of quicklime) at a temperature not exceeding 40° C and kept from access of air in a stoppered bottle, the glass stopper being smeared with Vaseline, or they will be destroyed by mites and undergo partial decomposition. The *Mylabris phalerata*, Pall., which is given in the *Ind* and *Col Add* as a permissible substitute for Cantharis in India and the African and Eastern Colonies, has elytra or wing cases, which are black with two wavy or transverse bands, and a large spot of the same colour at the base of each elytron. It is oblong, 25 mm or more in length and 9 mm in breadth. Other species containing 'the same percentage' of Cantharidin as *M. phalerata* (amount not stated) may be also employed for official purposes in those countries. Under the name of Chinese Cantharides it is occasionally found in English commerce, mixed largely with the smaller species, *Cichorus*, Fabr., these insects forming a cheap source of Cantharidin, but the *M. Cichorus* contains less than *M. phalerata*. *M. bifasciata*, Oliv., a Cape of Good Hope insect yields about 1 p.c. of Cantharidin. The new crop of Cantharides can be procured in early autumn. The *U S P* requires that Cantharides should be thoroughly dried at a temperature not exceeding 40° C (104° F) and that the powder should contain few or no hairs, but the insect itself is hairy. The microscopical characters of powdered Cantharides are given in *P J.* (4) xxv. p. 185.

Tests.—Numerous processes have been devised for the determination of Cantharidin. That adopted by the *P G.* is essentially as

follows —A weighed quantity of 25 grammes of the powdered Cantharides is treated with 100 grammes of Chloroform and 2 c c of Hydrochloric Acid, the mixture is allowed to stand for 24 hours with occasional agitation, and 52 grammes of the Chloroform solution is filtered off through a dry, closely-covered filter. The Chloroform is distilled off and the residue is treated with 5 c c of Petroleum Ether, allowed to stand for 12 hours with occasional shaking, the liquid filtered through a 5 cm filter paper, which has previously been weighed after extraction with Petroleum Ether, the residue is washed on to the filter with 10 c c of Petroleum Ether without endeavouring to detach the crystalline residue adhering to the flask. Both the flask and the filter are washed with Water containing a drop or two of Ammonium Carbonate Solution (1 of Ammonium Carbonate in 3 by weight of Water and 1 part by weight of Ammonia Solution) till the washings are colourless, then washed with 5 c c of Water and weighed. The crystalline residue should amount at least to 0.1 gramme corresponding to 0.8 p.c. w/w of Cantharidin.

This standard is considered (*P J* '01, n 715) to be too high, the average yield of commercial Spanish flies is nearer 0.6 p.c.

Myiabras Cichorni is richer in Cantharidin than Cantharides, and the use of these flies as a source of the vesicant has therefore been suggested. The employment of Cantharidin in place of Cantharides for the production of the galenical preparations has been advocated in several quarters. If this view were supported by clinical evidence it would obviate the difficulties encountered by the variations of commercial samples, and would enable other species of Cantharis possibly containing a larger percentage of the active principle to be utilised. The new Belgian Pharmacopœia employs Cantharidin for the preparation of an ointment.

The ash of Cantharides should not exceed 8.0 p.c.

Preparations

ACETUM CANTHARIDIS VINEGAR OF CANTHARIDES

Cantharides, bruised, 2, Glacial Acetic Acid and Distilled Water, mixed in equal volumes, *q s* to yield 20, by maceration for 24 hours and subsequent percolation (1 in 10)

A corresponding preparation, *Acetum Myiabridis*, is official in the *Ind* and *Col Add* for India and the African and Eastern Colonies.

It has been suggested (*P J* '98, 1 259, *C D* '98, 1 422) to replace Cantharides by Cantharidin, dissolving 1 of Cantharidin in 200 of Glacial Acetic Acid and adding Acetic Acid *q s* to yield 2000. This formula has been incorporated in *B P C* under the title *Acetum Cantharidini*.

Foreign Pharmacopœias—Official in *Port*, about 1 in 6, *Dutch* (*Acetum Myiabridis*), 1 in 10. Not in the others.

Tests—Vinegar of Cantharides has a specific gravity of 1.064 to 1.070, should contain not less than 2.5 p.c. w/v of total solids, and 1 c c should require for neutralisation not less than 8 c c of Normal Volumetric Potassium or Sodium Hydroxide Solution, indicating not less than 47.66 p.c. w/v of absolute Acetic Acid.

COLLODIUM VESICANS. BLISTERING COLLODION

Blistering Liquid, 20, Pyroxylin, $\frac{1}{2}$, dissolve by agitation in a well-closed vessel

U S is made with Flexible Collodion

See also CANTHARIDIN, below

Foreign Pharmacopœias—Official in Dan, Ger, Jap, Mex (Collodion Cantharidæd), Norw, Port, Russ, Swiss and U S Not in the others

EMPLASTRUM CALEFACIENS. B P Syn—WARMING PLASTER

Infuse 1 of Cantharides, in coarse powder, in 5 of boiling Distilled Water, strain and evaporate to $1\frac{1}{2}$ on a water-bath, add Yellow Beeswax, 1, Resin, 1, Resin Plaster, 13, Soap Plaster, 8

(about 1 in 25)

A corresponding preparation, **Emplastrum Calefaciens Mylabridis**, is official in the *Ind* and *Col Add* for India and the African and Eastern Colonies

EMPLASTRUM CANTHARIDIS. CANTHARIDES PLASTER

Cantharides, in powder, 7, Yellow Beeswax, 4, Lard, 4, Resin, 4, Soap Plaster, 1

(nearly 1 in 3)

Melt the Resin first, and add to it the plaster, Wax, and Lard, when all are completely fluid add the Cantharides gradually with stirring, continue the same until cold

A corresponding preparation, **Emplastrum Mylabridis**, is official in the *Ind* and *Col Add* for India and the African and Eastern Colonies

It has been suggested (*P J* '98, 1 259, *C D* '98, 1 422) to replace Cantharides by Cantharidin, dissolving 1 of Cantharidin in Chloroform *q s* and stirring it with 999 of a mixture in equal parts of Yellow Wax, Prepared Suet, and Resin The solution is stirred into the melted mixture and the Chloroform should be dissipated in the formation of the plaster This formula has been incorporated in *B P C* under the title **Emplastrum Cantharidin**

Foreign Pharmacopœias—Official in Belg, Dutch, Fr, Hung, Ital, Mex, Span and Swed, about 1 in 8, Austr, Dan, Ger, Jap, Norw, Port, Russ and Swiss, about 1 in 4 Not in U S

Emplastrum Cantharidum Perpetuum, Swiss, 3 in 10, Dan, Nor and Swed (*Emp Canth cum Euphorbio*), about 1 in 7, Hung, 1 in $5\frac{1}{2}$, Ital (*Empiastro de Cantaride mite*), 1 in 21, Austr, Ger and Russ, 1 in 10 Not in the others Norw includes an **Emplastrum Cantharidis Colatum**, Ger includes an **Emplastrum Cantharidum pro usu veterinario**, about 1 in 5

LIQUOR EPISPASTICUS. BLISTERING LIQUID

10 of Cantharides, percolated with Acetic Ether to produce 20 of Liquor

(1 in 2)

A corresponding preparation, **Liquor Epispasticus Mylabridis**, is official in the *Ind* and *Col Add* for India and the African and Eastern Colonies

Span has **Tinctura Chloroformica de Cantaridas**, White Wax, 1, Cantharides, 100, Chloroform, *q s* to make 100 By weight

See also CANTHARIDIN, below

TINCTURA CANTHARIDIS. TINCTURE OF CANTHARIDES

Macerate 1 of Cantharides, in No 40 powder, with 80 of Alcohol

(90 p c)

(1 in 80)

Dose.—5 to 15 minims = 0.3 to 0.9 c c ; if frequently repeated, 2 to 5 minims = 0.12 to 0.3 c c.

Ger and U S are much stronger

Ph Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes

It has been suggested (*P J* '98, 1 259, *C D* '98, 1 422) to replace Cantharides by Cantharidin, dissolving 1 of Cantharidin in 100 of Chloroform and adding Alcohol (90 p c) *q s* to yield 10,000

This formula has been incorporated in *B P C* with an increase of the Chloroform to 125, under the title **Tinctura Cantharidini**

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fl, Ger, Ital, Jap, Port, Russ, Span, Swed, Swiss and U S, 1 in 10, Mex, 1 and 10, Hung, 1 and 5. All by weight except U S

The *Brussels Conference* adopted a strength of 10 p c for the Tincture, employing Alcohol (70 p c)

Tests—Tincture of Cantharides has a specific gravity of about 0.835, it contains about 0.25 p c w/v of total solids and about 90 p c w/v of Absolute Alcohol

UNGUENTUM CANTHARIDIS CANTHARIDES OINTMENT

Cantharides, bruised, 1, Benzoated Lard, 10 digest at 120° F (48.9° C) for 12 hours and strain through calico, using gentle pressure towards the end (about 1 in 10)

Employed to promote discharge from a blistered surface. Being very painful this is seldom practised

A corresponding preparation, **Unguentum Mylabridis**, is official in the *Ind* and *Col Add* for India and the African and Eastern Colonies

It has been suggested (*P J* '98, 1 259, *C D* '98, 1 422) to replace Cantharides by Cantharidin, dissolving 1 of Cantharidin in Chloroform *q s*, and stirring it into a mixture of 499 of Yellow Beeswax and 2500 Olive Oil (by weight) previously melted. The Chloroform should be dissipated in the formation of the Ointment.

This formula has been incorporated in *B P C* under the title **Unguentum Cantharidini**, in which the proportions are given as 0.0325, 16, and 84

Foreign Pharmacopœias—Official in Fr (*Pommade Epispastique Verte*), about 1 in 83, and *P E Jaune*, 1 in 17½, Ger and Jap, Oil of Cantharides 3, Yellow Wax 2, Port, about 1 in 23, Ital (*Pomata di Cantharidi*), 1 in 10, Swed, 1 in 5 (fort, 1 in 4), Span, 3 in 10, Swiss, Cantharidine, 1 in 250, U S (*Ceratum Cantharidis*), 32 in 100, Mex (*Unguento de Cantharidas*), about 1 in 18. Not in Austr, Dutch or Hung. Ger and Swiss have **Unguentum Cantharidum pro usu veterinario** (1 in 5), Belg (**Unguentum Cantharidini**, *Cantharidin*), 1 in 2000, (**Unguentum Cantharidis cum Euphorbio**), *Cantharides*, 1 in 5

Ger and Jap have *Oleum Cantharidatum*, Mex has *Aceto de Cantharidas*

Not Official

CANTHARIDIN $C_{10}H_{12}O_4$, eq 194.62—White, inodorous, crystalline scales

Solubility—1 in 1150 of Rectified Spirit, 1 in 700 of Rectified Ether, sp gr 0.720, 1 in 55 of Chloroform, 1 in 150 of Acetic Ether, but even when dissolved at 60° F part separates on standing, 1 in 200 of Almond Oil, 1 in 65 of Oil of Cloves

Acetone is the best solvent for Cantharidin, which it dissolves 1 in 40, and as it is cheaper it possesses a double advantage over Acetic Ether. Acetone makes a good *Liquor Epispasticus*, it also dissolves Pyroxylin, and is therefore suitable for making *Collodium Vesicans*. Acting upon this suggestion *B P C* have included a **Collodium Cantharidini**—Cantharidin, 0.35 gramme, Acetone Collodion, *q s* to make 100 c c, but in the *B P C Supplement* the Cantharidin is first dissolved in 30 c c of Acetone, then Acetone Collodion added to make up 100 c c

Foreign Pharmacopœias,—Official in Belg, Fl, Mex, Port, Span and Swiss

Tests—Cantharidin melts at 218° C (424 4° F), and when heated further sublimes in white needles. The aqueous solution is neutral towards Litmus paper. Solutions of Cantharidin possess powerful vesicating properties, it dissolves without change of colour in concentrated Sulphuric Acid, and again separates when diluted with Water. It leaves no residue upon ignition.

POTASSIUM CANTHARIDATE—Colourless, needle-shaped crystals, soluble in Water, insoluble in Ether and in Chloroform. Should be preserved in well-stoppered bottles.

Liebreich's Solution contains 0.2 gramme Cantharidin and 0.4 gramme Potassium Hydroxide in 1000 c.c. of sterilised Water, 1 c.c. contains 0.0002 gramme Cantharidin in the form of Potassium Cantharidate. **Dose**—0.5 c.c., given also internally in diseases of tubercular origin, in lupus, and in _____ once been replaced by a tincture made with Cantharidin, 1, in Tincture of Orange Peel, 5000. **Dose**—0.5 c.c., and never more than 0.75 c.c., mixed with liqueur-glassful of Water.—*B M J* '02, ii 1231, *P J* '02, ii 708.

Official in Fr, Mex. and Span.

CHARTA EPISPASTICA (*B P* 1885)—Powdered Cantharides, 4, White Wax, 16, Spermaceti, 6, Olive Oil, 8; Resin, 3, Canada Balsam, 1, Distilled Water, 24. Conveniently spread on paper ruled in divisions of 1 square inch. Fr has Sparadiap Vesicant and Sparadiap de Cantharidate de Potassium.

EMPLASTRUM VESICANS—Cantharidin, 1, Chloroform, a sufficiency, Yellow Beeswax and Wool Fat, in equal proportions, 499 parts. The Chloroform is used to dissolve the Cantharidin, and is afterwards dissipated on a water bath.—*University*.

LINIMENTUM CRINALE—Cantharidin, 1 grain, Acetic Ether, 6 fl. drms., dissolved in 6 fl. oz. add Alcohol (90 p.c.), 6 fl. oz., Castor Oil, 2 fl. oz., Oil of Lavender, 15 minims.

This Liniment is highly recommended for application to the head where the hair is falling off, but after applying it a few times the head should be washed or the Cantharidin may accumulate, and cause too much irritation. It may be diluted with equal parts (or more) of Alcohol (90 p.c.) for delicate skins.

LIQUOR CANTHARIDIS CONCENTRATUS—1 fl. oz. = 1 oz. of Cantharides. It is obtained by re-percolation with Acetic Ether, and is standardised to contain 0.5 p.c. of Cantharidin. This Liquor forms a convenient substitute for Cantharides in making the various preparations, it effects a great saving of time and produces a better result.

Acetone is better as a solvent, but cannot at present be employed for official preparations.

UNGUENTUM STIMULANS (Erasmus Wilson's)—Cantharides, in Powder, 3 Lard, 12, macerate with a moderate heat for twenty-four hours, and filter through paper.

In place of the Cantharides, 6 of *L. _____* 3 of Liquor Cantharidis Concentratus may be employed, _____ extract, and mixed with the same on Lard.

BONI'S BLISTER—Camphor, 20, Chloral Hydrate, 30, melt and add powdered Cantharides, 10, digest for an hour at 150° F, filter.

CAOUTCHOUC.

INDIA-RUBBER

Fr, CAOUTCHOUC, Ger, KAUTSCHUK, Ital, CAUCCIÙ, Span, CAUCHO.

The prepared milk-juice of *Hevea brasiliensis*, and various other species. The best commercial variety is known as Para rubber.

Official Preparation—Tincture of Caoutchouc. The Liquor is used in the preparation of Chloroform emulsion.

Foreign Pharmacopœias—Fr, Ger, Jap, Mex, Span, Swiss and U S (Elastica) Not in the others

Descriptive Notes—In Brazil Para rubber is obtained from *Hevea brasiliensis*, Muell Arg, and other species, but in Ceylon from *Hevea brasiliensis* only

The *BP* statement that it is brownish black externally and mottled with a pale tint internally, except that mottling scarcely describes the gradual paling from the surface to the centre, applies also to Ceylon or biscuit Para rubber, this is now imported and is a remarkably pure product, though deficient in the antiseptic bodies and empyreumatic odour derived from the smoke of burnt Palm nuts that characterises the Brazilian Para rubber. Caoutchouc is easily cut with a knife if first wetted with Water

Tests—Caoutchouc melts at about 125° C (257° F), which figure is given in both *USP* and the *BP*, the *PG* gives 120° C (248° F). When melted it forms a fluid or semi-fluid mass, which on strongly cooling again becomes solid and still maintains its stickiness. It is insoluble in Water, Alcohol (90 p c) and in dilute solutions of the alkali Hydroxides. It dissolves in Benzol, Carbon Bisulphide, Chloroform, Petroleum, Ether and Oil of Turpentine. This statement must not be taken to mean that the rubber is entirely soluble in these liquids, but when treated with them it swells up and becomes gelatinous and soft, a portion being left in a disintegrated condition. This behaviour would suggest that rubber consists of constituent parts, one constituent part of which dissolves leaving the less soluble part in an insoluble but disintegrated condition

Preparation

LIQUOR CAOUTCHOUC SOLUTION OF INDIA-RUBBER

India rubber, 1, Benzol, 10, Carbon Bisulphide, 10

CAPSICI FRUCTUS

CAPSICUM

FR POIVRE DE GUINÉE, GFR, SPANISCHER PFEFFER, ITAL, PEPPERON, SPAN, PIMIENTO DE INDIAS

The dried ripe Fruit of *Capsicum minimum*

Imported from Zanzibar, Sierra Leone, etc, and distinguished in commerce as Guinea Pepper, Chillies, or Bird Pepper. That from Nepal has the finest flavour, and the powdered fruit is often preferred to the ordinary Cayenne Pepper

It yields its virtues to Water, Alcohol, Ether, Acetic Ether, and the fixed and volatile Oils

Medicinal Properties—Stomachic and carminative, used chiefly as a condiment. Given in dyspepsia and flatulent distension, and to promote appetite in alcoholism. Used externally as a rubefacient, and counter-irritant in rheumatism and lumbago and for chilblains

Dose.—½ to 1 gram = 0.01 to 0.06 gramme in pill

Tinct Capsici, 1½ dm (increased), Tinct Aurant, 4 dm. Syr Aurant, 4 dm Quinine Hydrochloride, 6 grams, Water, to 6 oz Take a tablespoonful as required, three to four times a day, in dipsomania

Official Preparations—Tinctura Capsici, and Unguentum Capsici The Tincture is contained in Tinctura Chloroformi et Morphinæ Composita

Not Official—Emplastrum Capsici, Extractum Capsici Liquidum, Gossypium Capsici, Linimentum Capsici, Liquor Capsici Compositus, Oleo-resina Capsici, Tinctura Capsici Ætherea, Tinctura Capsici Fortior, Unguentum Oleo-resina Capsici, and Capsicum with Wool Fat

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Dutch, Ger, Jap, Mex (Chile), Port (Pimenta o), Russ, Swed, Swiss and U S Not in the others

Descriptive Notes—The fruits of *Capsicum minimum*, Roxb, which are officially called Capsicum, are known in commerce as Chillies or Bud Pepper, they vary considerably in pungency and colour Those of Sierra Leone are usually yellowish-red, without pedicel or calyx, and are the most pungent of all Those from Zanzibar are redder and usually have the stalk and calyx attached, and are somewhat less pungent The Japanese are bright red, much less pungent, and the larger variety exceeds the dimensions given in the B P Like all ordinary Capsicum fruits the pericarp is glabrous, translucent and coriaceous, the seeds are flat and 10 to 20 in number The official description limits it to those having the following characters—Colour, dull orange-red, shape, ' ' and obtuse, size, ½ to ¾ in (12 to 20 mm) long, and ¼ in (6 mm) in diameter, the calyx and slender peduncle may be present or not The bright red Cayenne Pepper of commerce is usually prepared from Natal or Egyptian varieties of *Capsicum annuum*, L, which are eight or nine times as large as the fruit of *Capsicum minimum* Nepal Cayenne Pepper is usually of yellowish-brown tint, and has an odour of Violets, and is prepared from the yellowish-red fruit of the Nepal variety of ' ' The bright red Bud Pepper that is given to canaries, and has hardly any pungency, is prepared from the fruit of *Capsicum annuum* var *grossum*, Sendtn, grown in Spain, and known there as 'Pimento' or from the fruit of *Capsicum tetragonum*, Miller, grown in Hungary, and known there as 'Paprika' In both countries the fresh fruit is used as a condiment with food Coconada Capsicums are said to be the produce of *Capsicum annuum*, L, var *abbreviatum*, Fingerh and Natal and Indian Capsicums of the var *acuminatum*, Fingerh The bright red Cayenne Pepper of commerce is largely made from the variety imported from Natal J. E Wallis has shown that the powder of *Capsicum minimum*, *Capsicum annuum*, and Japanese Chillies can be distinguished under the microscope and suggests that the following description should be included in the Pharmacopœia as a means of excluding substitutes for, or admixtures with, the official Capsicum—'The pericarp shows an epidermis of thick and ' ' cells which have few pits, are often arranged in groups of 5 or 7 in a row and have an evenly striated cuticle' (See Pharm Jour (4) xiii p 552, xv p. 3) The U S P requires that the powdered Capsicum contains few or no starch grains or sclerenchymatous fibres and refers the matter to C. C.

fastigiatum, Blume The large fruits of *Capsicum annuum*, 5 to 10 cm long, and 4 cm at the base, are official in the *P G* According to Geriand, Alcohol (90 p c) is the best and most perfect solvent of the active principle of Capsicum (*Y B P* '05, 453, *P J* (4) xxi p 153) The pungent principle is most abundant in the placenta which yields 0.9 p c, the rest of the fruit yielding only 0.2 p c The powder of Capsicums soon becomes mouldy, if not kept dry

Tests—Capsicum leaves about 6.0 p c of ash, which figure should not be exceeded

Preparations

TINCTURA CAPSICI TINCTURE OF CAPSICUM

Macerate 1 of Capsicum, in No 20 powder, with 20 of Alcohol (70 p c) (1 in 20)

Dose—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Belg, 1 and 10, Mex, 1 in 5, Dan and Dutch, 1 and 10, Ger, Jap, Russ, Swed and Swiss, 1 in 10, all by weight U S, 1 in 10 Not in the others

Tests—Tincture of Capsicum has a specific gravity of about 0.895, contains from 0.7 to 1.5 p c w/v of total solids and about 70 p c w/v of Absolute Alcohol

UNGUENTUM CAPSICI CAPSICUM OINTMENT

Bruised Capsicum, 120 grains, Spermaceti, 60 grains, Olive Oil (by weight), 1 oz, strain after digestion on a water-bath for one hour

It has been suggested to use half the quantity of Liquid Extract of Capsicum (2 in 1) in place of Capsicum

Not in the Foreign Pharmacopœias

Not Official

CAPSICUM WITH WOOL FAT—1 of Liquid Extract of Capsicum (2 in 1), incorporated with 9 of Hydrous Wool Fat This was suggested by Geriand as an improvement on the ointment

EMPLASTRUM CAPSICI (*Geriand*)—Liquid Extract of Capsicum (2 in 1), 10, Resin Plaster, 95, evaporate the Alcohol and mix the residue with the plaster This formula closely resembles that incorporated in *B P C*

EMPLASTRUM CAPSICI (*U S*)—Apply a thin coating of Oleo resin of Capsicum, by means of a brush, so as to form a thin coating over an area 15 centimetres square, leaving a margin round the sides

EXTRACTUM CAPSICI LIQUIDUM (*Geriand*)—Exhaust 100 of Capsicum Fruit in No 60 powder with Alcohol (90 p c), distil off the Alcohol until the needed extract weighs 50 This has been incorporated in *B P C* Fluid extractum Capsici (*U S P*) is half this strength

GOSSYPIUM CAPSICI—Geriand's form is to saturate evenly 9 of Cotton Wool under pressure, with a mixture of 2 of Liquid Extract of Capsicum (2 in 1) and 7 of Alcohol (90 p c), and then dry It is coloured with Eosin to keep the colour more uniform *B P C* formula is just half the strength of this

LINIMENTUM CAPSICI—Stronger Tincture of Capsicum, 35 c c, Oleic Acid, 12.5 c c, Oil of Lavender, 0.625 c c, Alcohol, q s to make 100 c c This preparation corresponds to Linimentum Capsici, *B P C*, and to the Liniment recommended in Martindale, 1906

Oak, Beech, Hazel, Willow, and Poplar are employed

Medicinal Properties—Antiseptic, absorbent and deodoriser Given in cases of distension by intestinal gas, and in foul eructations and diarrhoea in dysentery and typhoid, also in dyspepsia attended with flatus, acidity and pain It will absorb and neutralise poisonous alkaloids Externally, as a poultice, it cleanses and absorbs the fetor of ulcers and gangrenous parts

Dose—60 to 120 grains = 3.9 to 7.7 grammes

Prescribing Notes—*It has been given in powder diffused in Water, also in the form of capsules, cachets, and biscuits The most palatable way is to mix it with chocolate*

Foreign Pharmacopœias—Official in all except Dan, Jap, Norw and Swed, Mex (Carbon Vegetal)

Tests—It should, according to the official requirements, leave 7.5 p.c. of ash when burned at a high temperature with free access of air, the *P G* permits only an insignificant amount of ash, and that it should burn without flame The *USP* includes a test with Potassium Hydroxide Solution which ensures evidence of complete carbonisation

Potassium Hydroxide—If 1 gramme be boiled with a mixture of 3 c.c. of Potassium Hydroxide T.S. and 5 c.c. of Water for several minutes, the filtrate should be colourless or nearly so, *USP*

CARBONIS BISULPHIDUM.

CARBON BISULPHIDE

B P Syn—CARBON DISULPHIDE

CS_2 , eq 75.55

A colourless, very volatile, highly refractive, limpid liquid, having an ethereal and not unpleasant odour when quite pure, but usually possessing a very disagreeable odour due to impurity

Should be preserved in well-stoppered dark amber-tinted glass bottles, partially filled, or in tin cans, in a cool atmosphere and away from naked flames, and not exposed to light

It is produced by heating Charcoal and Sulphur together at a high temperature, the crude product being condensed and subsequently rectified

Solubility—About 1 in 500 of Water, readily soluble in Absolute Alcohol, Ether (sp. gr. 0.720), Chloroform, and the fixed and volatile Oils

It is a good solvent for Iodine, Phosphorus, Precipitated Sulphur, etc

Medicinal Properties—It is official as a solvent for India-rubber and Phosphorus It is a powerful poison, and is not often given internally

1 or 2 oz. daily of a saturated Solution in Peppermint Water have been given as a substitute for Beigeon's treatment of phthisis—*B M J* '88, 1.421

A claim has been made that this should not remain in obscurity, but retain its place as a valuable remedy in tuberculosis—*B M J E* '04, 11.72.

Internally in pneumonia, in 5 p c aqueous solution hourly -*B M J E* 06,
11 68

Official Preparations—Used in the preparation of Liquor Caoutchouc and Pilula Phosphori

Foreign Pharmacopœias—Official in Belg, Fl, Port, Spain and U S Not in

Tests—Carbon Bisulphide has a specific gravity of about 1.268, *USP* gives 1.256 to 1.257 at 25° C (77° F), and a boiling point of about 46° C (114.8° F), the *BP* and *USP* give 46° to 47° C (114.8° to 116.6° F). It burns with a bluish flame, yielding Carbon Dioxide and Sulphur Dioxide as products of combustion.

The more generally occurring impurities are dissolved Sulphur, Sulphur Dioxide, and Hydrogen Sulphide. Dissolved Sulphur is shown by the residue left on evaporation, Sulphur Dioxide by its bleaching action towards moistened blue Litmus paper, and Hydrogen Sulphide by Lead Acetate.

Litmus—Blue Litmus paper moistened with Water should not be affected by Carbon Bisulphide, *BP* and *USP*.

Residue.—When a portion of Carbon Bisulphide is allowed to evaporate spontaneously in a glass vessel, no residue should be left, *BP* and *USP*.

Lead Acetate—Lead Acetate T.S., agitated with Carbon Bisulphide, should not be blackened, *BP* and *USP*.

CARDAMOMI SEMINA.

CARDAMOM SEEDS

FR, CARDAMOMES, GER, KARDAMOMEN, ITAL, CARDAMOMO,
SPAN, CARDAMOMO MENOR

The dried, ripened Seeds of *Elettaria Cardamomum*. *BP* states that the seeds should be kept in their pericarps, and separated when required for use.

1 of Fruit yields about $\frac{1}{3}$ of Seeds

Medicinal Properties.—Stomachic, carminative, and flavouring agent, a useful adjuvant to papaveres to prevent griping.

Official Preparation.—Tinctura Cardamomi Composita. Contained in Extractum Colocynthis Compositum, Pulvis Cinnamomi Compositus, Pulvis Cretæ Aromaticus, Tinctura Gentianæ Composita, Tinctura Rhei Composita. Of the Tincture contained in Decoctum Aloes Compositum, and Mistura Sennæ Composita.

Not Official—Oleum Cardamomi, Tinctura Cardamomi, Tinctura Carminativa, and Mistura Carminativa.

Descriptive Notes.—The Cardamoms of commerce are derived from several species, but the official kind is limited to the fruits of typical *Elettaria Cardamomum*, Maton, by the measurements given, viz, $\frac{3}{4}$ to $\frac{1}{2}$ in (1 to 2 cm), and by the 'pale buff' colour. The varieties of this Cardamom, as met with in trade, are known as Malabar, Mysore, and Mangalore, whether imported from those districts or from Ceylon, where Cardamoms are largely cultivated. The Malabar variety consists of short, plump, firm capsules well filled

with seeds, those of Mangalore are similar, but more or less watery on the surface, and those of Mysore are longer and less compactly filled with seeds, and consequently can be more easily compressed between the fingers. These varieties may also be met with in a bleached form, obtained by moistening them and submitting them to the action of Sulphurous Acid gas, in which case they present a smoother surface, the natural longitudinal striations being somewhat obscured in the process. For galenical purposes the shorter fruits showing the natural striations are to be preferred since their natural colour shows their good quality and their plumpness indicates a large proportion of seed to husk, and seeds more fully matured than in the Mysore kind. The fruits are usually collected before they are quite ripe to prevent the pericarp splitting open. Such partially open fruits as do occur are apparently sorted out and husked by being passed between rollers, or by similar means, since a certain amount of 'split' seed is offered in commerce.

The seeds should not be removed from the pericarps until required for use. The loose seeds obtainable in commerce present the possibility of being obtained from other than the official species, and in any case are likely to be deficient in aroma from exposure to the air.

The distinguishing microscopical characters of the powdered official Cardamoms are the perisperm cells, containing small starch grains, and prismatic Calcium Oxalate crystals, the dark coloured polyhedral cells of the inner integument, and the thick walled linear cells with oblique ends, of the epidermal layer. The presence of the pericarp in the powder may be detected by the straight-walled, polygonal cells of the epidermal parenchyma, spiral vessels, and small cells containing brown resin.

The other varieties of Cardamom occurring in commerce at more or less regular intervals are the var *major* of the official kind, known as Wild Ceylon Cardamom, which have longer, greyish fruits, the Greater or Korarima Cardamom (*Amomum Korarima*, Perena), which is about $1\frac{1}{2}$ in (37 mm) long and $\frac{1}{4}$ in (19 mm) broad at the base, and of a dull brownish colour, the cluster Cardamom (*Amomum Cardamomum*, L.), which is whitish, spherical, nearly smooth, and about $\frac{1}{2}$ in (12.5 mm) in diameter, all of these have seeds resembling the true Cardamom in flavour. The Bengal Cardamom (*A. aromaticum*, Roxb), the Nepaul (*A. subulatum*, Roxb) and the bitter-seeded Cardamom, all having a brown pericarp, are more rarely imported, but the seeds, freed from the husk, and when offered in commerce in the form of powder for use in pills are not so easy of recognition as the fruit, and are best detected by a microscopical examination, see *PJ* (4) vi p 280. The official seeds are dark reddish-brown and 3 mm ($\frac{1}{8}$ in) in diameter, angular and transversely wrinkled.

Tests—Cardamoms are officially required to yield not more than 4 p.c. of ash. Determinations of the ash of Pericarps, Seeds, and Pulvis made in the author's laboratory yielded Pericarps (three samples), 10.4, 12.0, 13.4 p.c., Seeds (three samples), 2.38, 2.81, 3.85 p.c., Pulvis (three samples), 7.56, 6.33, 9.93 p.c., these results

seem to indicate that the Pulvis Cardamomi was not obtained from the seeds only as directed in the Pharmacopœia. Even whole fruits had but an average of 5.5 p c. A maximum ash limit of 6 p c has been recommended.

Foreign Pharmacopœias.—The Fruit is official in Austri, Dan, Dutch, Ger, Hung, Jap, Mex (Cardamomo menor), Norw, Port, Russ, Swed, Swiss and U S. Not in Ital.

Preparation

TINCTURA (1875) COMPOSITA. COMPOUND TINCTURE OF CARDAMOMS

Cardamom Seeds, bruised, 1 oz, Caraway Fruit, bruised, 1 oz, Raisins of commerce, freed from seeds, 8 oz, Cinnamon Bark, bruised, 2 oz, Cochineal, in powder, 220 grains, macerated with 80 fl oz of Alcohol (60 p c) (1 in 80)

Dose.— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in U S, 1 in 40, contains Glycerin, and is made with the Fruit of the Cardamoms. Not in the others.

Tests.—Compound Tincture of Cardamoms has a sp. gr. of 0.945 to 0.950, it contains about 6 p c w/v of total solids and about 56 p c w/v of Absolute Alcohol.

Not Official.

OLEUM CARDAMOMI—A pale yellow aromatic oily liquid distilled from Cardamom Seeds, which contain about 4 to 6 p c.

Tests—Cardamom Oil is distilled chiefly from Ceylon Cardamoms. It has a specific gravity of 0.933 to 0.943, an optical rotation of $+26^{\circ}$ to $+34^{\circ}$, in a 100 mm tube. It is soluble in 4 parts and more of Alcohol (70 p c), and should possess a Saponification value of 132.

Schimmel states that Mysore (Ceylon) Cardamom Oil has a specific gravity of 0.895 to 0.905, but Parry has been unable to confirm these figures and shows that there is practically no difference between the two Oils.

TINCTURA CARDAMOMI—Cardamom Seeds, bruised, 1, Alcohol (60 p c), $q\ s$ to yield 10, by percolation.

Dose.—30 to 60 minims = 1.8 to 3.6 c c

The B P C Tincture, also 1 in 10, is made by maceration.

Foreign Pharmacopœias—Official in U S, 1 of Fruits in 5 of Alcohol (48.9 p c), by percolation, Port and Swiss, 1 in 5 (weight).

TINCTURA CARMINATIVA (B P C Formulary 1901)—Cardamom Seeds, bruised, 600 grains, stronger Tincture of Ginger (B P '85), 1 $\frac{1}{2}$ oz, Oil of Cinnamon, 100 minims, Oil of Caraway, 100 minims, Oil of Cloves, 100 minims, Alcohol (90 p c), $q\ s$ to yield 20 fl oz, macerate the Cardamoms in 15 fl oz of the Spirit for a week, decant, express, and dissolve the Oils in the mixed Tinctures, and add the remainder of the Alcohol.

Dose.—2 to 10 minims = 0.12 to 0.6 c c

This has been incorporated in the B P C.

By replacing the Cardamom Seeds with $\frac{1}{8}$ of Oil of Cardamoms the maceration is avoided.

MISTURA CARMINATIVA—Sodium Bicarbonate, 60 grains, Aromatic Spirit of Ammonia, 72 minims, Compound Tincture of Cardamoms, 144 minims, Glycerin, 240 minims, Dill Water, to 6 $\frac{1}{2}$ fl oz.

This corresponds to Mistura Carminativa (B P C).

Several formulas are given in Ph. Form.

CARUI FRUCTUS.

CARAWAY FRUIT

FR, CARVI, GER, KUMMEL, ITAL, CARVI, SPAN, ALCARAVEA

The dried Fruit of *Carum Carvi*, L

Cultivated in different parts of Europe The herb flowers in the second year, and the fruit ripens in July or August Yields from 8 to 7 p c of Oil, varying with the source of the Seeds

Medicinal Properties—Aromatic, stomachic, and carminative Used in flatulent colic, as an adjuvant to other medicines, to prevent griping of purgatives, and as a flavouring agent

Official Preparations—Aqua Carui, and Oleum Carui Contained in Confectio Piperis, Pulvis Opii Compositus, Tinctura Cardamomi Composita, Tinctura Sennae Composita The Oil is contained in Pilula Aloes Barbadosis

Foreign Pharmacopœias—Official in Austr, Ger, Mex (Alcaravea), Port (Alcaravie), Swed, Swiss and U S Not in the others

Descriptive Notes—The principal varieties of Caraway fruit met with in commerce in this country are the English, Dutch and Russian, which differ in size, colour and aroma The English command the highest price, and are light brown in colour and slightly larger than the Dutch, which are a dark brown and cheaper, and are the kind usually sold by grocers According to the official description they should be 'about' $\frac{1}{4}$ to $\frac{1}{2}$ in (4 to 6 mm) long and about $\frac{1}{8}$ in (1 mm) broad, tapering towards each end, and would include these two The Russian Caraways are small and mixed with a considerable quantity of stalks and debris and are chiefly used in veterinary medicine The Mogador Caraway is occasionally imported It is the largest of all, and is usually slightly enlarged at the upper end, and consequently would be excluded by the *B P* statement that they are tapering at each end The Dutch, Norwegian and East Russian are chiefly used in Germany for distillation The value of the oil depends upon the amount of Carvone it contains The exhausted seeds are dried and sometimes used for purposes of adulteration, but they may be detected by their darker colour, weaker taste, and shrivelled appearance, and (under a good lens) by the torn outer layer of cells Under the microscope the most noticeable features of the powder are the pitted walls of the cells of the outer epidermis, which, like those of Anise, are striated, but are more oblong in outline, and about twice as long as broad, the parallel, thin walled, elongated, oblong cells of the inner epidermis about four times as long as broad, the absence of raphides and hairs

Tests—Caraway Fruit leaves about 6 p c of ash on incineration, and 8 p c should not be exceeded Six samples of the Fruit examined in the author's laboratory showed from 5.72 to 7.1 p c, 5 samples of the powdered fruit gave from 5.87 to 7.15 p c

Preparations

AQUA CARUI. CARAWAY WATER

Caraway Fruit, 1, Water, 20, distil, 10.

(1 in 10)

Dose.—1 to 2 fl oz = 28.4 to 56.8 cc

Foreign Pharmacopœias—Official in Jap and Swed Not in the others

OLEUM CARUI. OIL OF CARAWAY

A colourless or pale yellow, mobile liquid, possessing a characteristic and aromatic odour and a spicy taste

The Oil distilled from Caraway Fruit It consists principally of a terpene, Dextro-limonene, and an oxygenated compound of a ketone nature, Carvone It is the latter to which the oil owes its medicinal properties

It should be kept in well-stoppered amber-coloured glass bottles and protected as much as possible from the light It should be kept in a cool place

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 cc

Foreign Pharmacopœias—Official in Austri (Carvone), Ger, Jap (Carvonum), Port, Swiss and US, Swed (Carvone) Not in the others

Tests—Caraway Oil has a specific gravity of 0.907 to 0.920 and an optical rotation of from $+70^{\circ}$ to $+80^{\circ}$ in a 100 mm tube The *BP* gives the specific gravity but no optical rotation, neither does it make any mention of the solubility of the oil in Alcohol (90 p c), it gives the specific gravity as 0.910 to 0.920, the *USP* gives 0.900 to 0.910 at 25°C (77°F) The *USP* gives the optical rotation as $+70^{\circ}$ to $+80^{\circ}$ in 100 mm tube at a temperature of 25°C (77°F) The oil should yield a clear mixture with an equal volume of Alcohol (90 p c), and with 3 to 10 volumes of Alcohol (80 p c) The *USP* gives the Alcohol solubility As pointed out above, Carvone is official in place of the oil in several Foreign Pharmacopœias, and commercial oils are frequently met with from which the Carvone has been abstracted Such abstraction is shown by the specific gravity and by the optical rotation, oils from which the Carvone has been taken having a specific gravity of about 0.848 and an optical rotation of over $+100^{\circ}$

Carvone forms crystalline compounds with Hydroxylamine, with Hydrogen Sulphide, and with Phenylhydrazine, but its actual determination, owing to difficulties inseparable from the process, can only be made with approximate accuracy A measured quantity of 5 cc of the oil is treated in a test-tube with 5 cc of Phenylhydrazine and the tube allowed to stand in boiling Water for an hour The excess of Phenylhydrazine is removed by adding whilst hot 5 cc of Glacial Acetic Acid, and after $\frac{1}{2}$ hr Water to 20 cc The contents are then cooled, the crystals removed by filtration and washed with Water until of a pale yellow colour They may be recrystallised from a definite volume of Alcohol (90 p c)

Not more than 15 p c of the oil should distil below 185°C (365°F), and at least 55 to 65 p c should distil above 200°C (392°F), the fraction between 220° to 230°C (428 to 446°F) should amount to a

The higher the specific gravity and the greater the solubility in Alcohol (50 p c) the more Carvone is likely to be contained in the sample

Not Official

CARVONE—A colourless or pale yellow fluid possessing a characteristic aromatic odour and taste. When obtained from the Oils of Caraway and Dill it is stated to be dextrogyrate, and lævogyrate when obtained from Oil of Spearmint.

Tests—Carvone has a specific gravity of not less than 0.960 and a boiling point of 229° to 230° C (444.2° to 446° F). It should be soluble in 2 parts by weight of Alcohol (68 to 69 p c). It forms crystalline compounds with Hydroxylamine, Hydrogen Sulphide and Phenylhydrazine, and may be determined quantitatively by means of Hydroxylamine or Phenylhydrazine. Carvone which has been exposed to the air when dissolved in an equal volume of Alcohol (90 p c) yields a reddish violet coloration with Ferric Chloride Test solution, which disappears on the further addition of Ferric Chloride Test solution.

It is official in Austr., Ger., Jap. and Swed. in place of the Oil

CARYOPHYLLUM

CLOVES

FR., GIROFLE, GER., GEWURZNELKEN, ITAL., GAROFANI,
SPAN., CLAVO DE ESPECIA

The dried Flower-buds of *Eugenia caryophyllata*

Imported from Penang, Bencoolen, Amboyna, and Zanzibar. Yield from 15 to 18 p c of Oil

Medicinal Properties—Aromatic, stomachic, carminative, antispasmodic. Administered to check nausea, vomiting, and flatulence, and to promote digestion. But chiefly used as an adjuvant to other medicines. The oil, as a counter-irritant, is a useful ingredient in liniments for whooping-cough and bronchitis, it is also used as an anodyne in toothache.

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes—*The Oil may be given upon a lump of Sugar, and is a useful constituent of aperient pill masses. The Infusion is a nice flavouring for many mixtures.*

Incompatibles—See under Infusum Caryophylli

Official Preparations—Infusum Caryophylli, and Oleum Caryophylli. Used in the preparation of Infusum Aurantii Compositum. Contained in Pulvis Cistæ Aromaticæ. The Oil is contained in Pilula Colocynthis Composita, and Pilula Colocynthis et Hyoscyami.

Not Official—Infusum Caryophylli Concentratum and Eugenol

Foreign Pharmacopœias—Official in Austri., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex. (Clavo de Especia), Norw., Port. (Clavinho), Russ., Span., Swed., Swiss and U.S.

Descriptive Notes—Cloves consist of the dried flower-buds of *Eugenia caryophyllata*, Thunb., the lower portion of which is formed of a calyx tube, enclosing in its upper half the ovary filled with minute ovules. It derives its name from the French word for a nail, *clou*, from its resemblance to a short nail in shape. The finest varieties in English commerce are imported from Penang, Bencoolen and Amboyna, and in French commerce from Réunion and Madagascar.

Those from Ceylon and the Seychelles are of medium quality and size and darker coloured. Those from Zanzibar and Pemba are more slender and the globular head is often broken. These last two varieties form about four-fifths of the world's production. The official Cloves should be $\frac{3}{8}$ in (15 mm) long, and should emit oil when indented with the finger nail. The 'stems' or stalks of the flowers are imported separately, and used for distillation of oil. Cloves exhausted of oil by distillation have been used to adulterate cloves, but yield no oil when pressed with the nail and float when put in water. Powdered Cloves have been adulterated with clove stalks and with the fruit of Cloves, known in commerce as Mother Cloves. The former may be detected by the presence of sclerenchymatous cells and the latter by the presence of starch granules, neither of which occur in Cloves.

Tests—Cloves yield when incinerated about 5 p.c. of ash, and 8 p.c. should not be exceeded. Eight samples examined in the author's laboratory yielded from 4.78 to 5.4 p.c., 5 samples of the powder yielded from 5.2 to 6.97 p.c. of ash. The Ether extract should amount to 20 p.c. The B.P. limit of ash is 7 p.c., and the U.S.P. 8 p.c.

Preparations

INFUSUM CARYOPHYLLI. INFUSION OF CLOVES

Cloves, bruised, 1, Distilled Water, boiling, 40, infuse for fifteen minutes, strain (1 in 40)

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 c.c.

Incompatibles—Lime Water, salts of Iron, mineral acids, Gelatin

OLEUM CARYOPHYLLI. OIL OF CLOVES

FR, ESSENCE DE GIROFLE, GÉR, NFKENOL, ITAL, ESSENZA DI GAROFANI, SPAN, ESENCIA DE CLAVO

A pale yellow, limpid, highly refractive liquid possessing a characteristic aromatic odour and taste, distilled from Cloves. The yield of oil is from 15 to 18 p.c.

It becomes darker in colour with age and on exposure to air, and should therefore be kept in well-stoppered, dark amber-tinted glass bottles, and protected as far as possible from contact with the air.

The oil contains from 70 to 85 p.c. of Eugenol, a phenol having the formula $C_{10}H_{12}O_2$, a sesquiterpene, Caryophyllene, Methyl Alcohol, Furfural, and a trace of Vanillin.

Solubility—1 in 60 of Alcohol, (60 p.c.), in all proportions of Alcohol (90 p.c.), Ether, and Strong Acetic Acid.

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Foreign Pharmacopœias—Official in Austri (Eugenol), Belg. (Eugenol), Denm. (Eugenol), Fr, Ger (Eugenol), Hung. Ital. Jap, Mex, Norw. Port, Russ, Span, Swed (Eugenol), Switz and U.S.

Tests—Clove Oil has a specific gravity of 1.048 to 1.068, it is officially required to be not below 1.050, the U.S.P. gives 1.040 to

1.060 at 25°C (77°F). Some genuine oils are occasionally met with of a lower specific gravity than that given in the *BP*, but rarely lower than 1.045. Both the *BP* and the *USP* require the oil to form a semi-solid yellowish mass when shaken with an equal volume of strong Ammonia Solution, the *USP* also mentions that concentrated Potassium Hydroxide Solution produces a similar result. When dissolved in Alcohol (90 p.c.) it is officially required to yield a blue coloration on the addition of Ferric Chloride Test-solution. The *USP* dissolves 2 drops of the oil in 4 c.c. of Alcohol (94.4 p.c.) and adds a drop of Ferric Chloride Test-solution, when a bright green colour will be produced, but a drop of a solution of Ferric Chloride Test-solution diluted with 4 times its volume of Water produces a blue coloration changing to yellow. This is as far as the official tests take us.

The optical rotation of the oil is about -1° in a 100 mm tube. No appreciable quantity of the oil should distil below 246°C (474.8°F). Eugenol may be determined approximately by treating a weighed quantity of 10 grammes of the oil in a flask with a long graduated neck with 100 c.c. of a 10 p.c. Potassium Hydroxide Solution, adding sufficient of the Hydroxide Solution to bring the level of the aqueous liquid to the zero mark, and reading off the volume of the unabsorbed portion which rises to the surface. This volume multiplied by 0.908 (the specific gravity of Caryophyllene) gives approximately the percentage by weight of the latter. The results are only approximate, owing to the solubility of the Caryophyllene in the Potassium Hydroxide Solution and Potassium Eugenate. The *USP* describes this process, but works by volume and not by weight. Thus determined, the *USP* Oil of Cloves is required to contain at least 80 p.c. of Eugenol. No method of determination is given in the *PG*.

A more accurate process is that of Thoms, which consists in converting the Eugenol into Benzoyl-eugenol by means of Benzoyl Chloride. A weighed quantity of 5 grammes of the oil is treated in a beaker having a capacity of about 150 c.c., with 20 grammes of Sodium Hydrate solution (15 p.c.) and 6 grammes of Benzoyl Chloride. The mixture is well shaken until uniformly mixed. After cooling, 50 c.c. of Water are added, and the mixture heated until the crystalline mass has again become oily, and is again allowed to cool. The clear supernatant liquid is filtered off and the crystalline mass in the beaker is again treated with two successive quantities of 50 c.c. of Water. The moist Benzoyl-eugenol is treated with 25 c.c. of Alcohol (90 p.c. by weight) and heated on a water-bath until solution is effected. The beaker is removed from the water-bath and agitated until the Benzoyl-eugenol has separated in fine crystals. The mass is then cooled to 17°C (62.6°F), the crystalline precipitate transferred to a small weighed filter paper, the filtrate collected in a graduated cylinder and washed with Alcohol (90 p.c. by weight) until it measures 25 c.c. The filter and crystals are transferred to a weighing bottle, dried at 100°C (212°F) till constant in weight and then weighed. The solubility of pure Benzoyl-eugenol in Alcohol

(90 p c by weight) has been experimentally proved to be 0.55 gramme, and the latter weight should be added to the weight of the crystals obtained. 266.11 parts of Benzoyl-eugenol represent 162.86 parts by weight of Eugenol.

The quality of Clove Oil is reduced by the abstraction of a portion of Eugenol, the addition of oil from Clove stems, and the addition of Turpentine or Petroleum. The abstraction of Eugenol is shown at once by the altered physical characteristics of the oil as well as by a determination of the amount of this constituent. Turpentine or Petroleum is revealed by a low boiling point and by the solubility of the oil in Alcohol (90 p c). Clove Oil from the stems can only be satisfactorily detected by the difference in odour, though the presence of Acetyl-eugenol in Clove Oil and not in oil from the stems has been suggested as a means of distinction.

The *U S P* includes a test for Phenol with Ferric Chloride T S, requiring that no blue or violet coloration should be produced when the oil is shaken with 20 times its volume of hot Water, cooled, the excess of oil removed by filtration through a wet filter paper and the filtrate tested with a drop of Ferric Chloride Test-solution.

Determination—A measured quantity of 10 c c of the Oil is introduced into a flask with a long neck graduated in tenths, and 100 c c of Potassium Hydroxide Test-solution added, the mixture being shaken for five minutes. After complete separation of the liquids, sufficient of the Potassium Hydroxide Solution is added to raise the lower limit of the oily layer to the zero mark of the scale, and the volume of the residual liquid is read off. This should not amount to more than 2 c c, indicating the presence of at least 80 p c of Eugenol, *U S P*.

Not Official

INFUSUM CARYOPHYLLI CONCENTRATUM—1 of Cloves in No. 10 powder, macerated for seven days in $2\frac{1}{2}$ of Alcohol (20 p c), and subsequently percolated so as to make 5 of concentrated Infusion.

This corresponds to Infusum Caryophylli Concentratum, *B P C*, and closely resembles Inf. Caryoph. Conc. given in *Ph. Form.* It is intended for dilution, 1 of this and 7 of Distilled Water.

EUGENOL—A phenol, having the formula $C_{15}H_{11}O_2$, eq. 162.86, is the principal constituent of Clove Oil. It is a colourless, pale yellow, highly refractive liquid, possessing a powerfully aromatic odour and taste. It darkens on exposure to air and light, and should be preserved in well-closed, dark amber-tinted glass bottles, and kept as far as possible from contact with the air.

Tests—Eugenol has a specific gravity of 1.072 to 1.074. Its boiling point is 251° to 253° C (483° to 487° F). It dissolves with difficulty in Water, but is readily soluble in Alcohol (90 p c), Ether, and Glacial Acetic Acid. A clear liquid, readily becoming turbid on exposure to air, is formed when 1 gramme of Eugenol is mixed with 26 c c of Water and 4 c c of Sodium Hydroxide Solution (15 p c). A flocculent precipitate, partially adherent to the sides of the vessel, is produced when 5 drops of Eugenol are shaken with 10 c c of Lime Water. An alcoholic solution of Eugenol yields with Ferric Chloride Test-solution a blue coloration, and with diluted Ferric Chloride Test-solution (1 to 10) a blue coloration changing to greenish yellow. One part by weight of Eugenol should be soluble in two parts by weight of Alcohol (68 to 69 p c). Eugenol is converted into a crystalline body by means of Benzoyl Chloride, and this property may be utilised for its determination. The method is described under Oil of Cloves. Eugenol is official in place of Clove Oil in the new editions of the *Austrian*, *Belgian*, and *Dutch Pharmacopœias*. It was made official in the 10th edition of the *German Pharmacopœia*, which includes a test for the presence of Phenol.

Eugenoforn (Sodium Eugenol Carbinol)—Colourless foliaceous crystals, readily soluble in water, slightly soluble in alcohol (90 p c), insoluble in ether. Introduced as an intestinal and stomachic antiseptic.—*P J* '99, 11 40

Dose— $7\frac{1}{2}$ to 15 grains = 0.5 to 1 gramme, twice a day

CASCARA SAGRADA.

CASCARA SAGRADA

B P Syn—*RHAMNI PURSHIANI CORTEX*, SACRED BARK

FR, CASCARA SAGRADA, GER, SAGRADARINDE, AMERIKANISCHE KREUZDORNRIINDE, ITAL, CASCARA SAGRADA, SPAN, CASCARA SAGRADA

The dried Bark of *Rhamnus Purshianus*

Obtained from California, best collected in spring and early summer. Bark which has been gathered for two years is much preferred to the recently dried bark

Medicinal Properties—Tonic laxative. Acts principally on the large intestine. Indicated in obstinate and habitual constipation, especially of old or delicate persons, and in an atonic condition of the stomach and bowels, as in anæmia. It should not be given as a purgative, but in such a constant continuous manner that a normal condition will be brought about. It is better to give two small doses, say 20 minims of the liquid extract night and morning, than one large dose. The dose should be reduced gradually.

Prescribing Notes—Usually given in the form of Extract in Pills or Pilules, or one of the fluid preparations. The Extract is best made into Pills with the addition of one tenth of its weight of Gum Acacia in powder, and massed with Alcohol (90 p c). It is also advantageously combined with Extract of Belladonna, Extract of Nux Vomica, and Euonymin. Obtainable in the form of Compressed Tablets. Capsules may be had containing a very concentrated Fluid Extract, equivalent to 15 and 30 minims of the ordinary Fluid Extract, and other strengths as desired. In Mucuses and other fluid preparations it goes well with Aromatic Spirit of Ammonia and Spirit of Chloroform.

Elixir of Cascara (Kasak) is an agreeable and reliable preparation. See below.

Official Preparations—Extractum Cascaræ Sagradæ, Extractum Cascaræ Sagradæ Liquidum, and Syrupus Cascaræ Aromaticus.

Not Official—Capsules of Cascara, Elixir of Cascara, Extractum Cascaræ Liquidum Insuperum, Mistura Cascaræ Sagradæ, Mistura Cascara Sagrada Composita, Mistura Cascara Aperiens, Mistura Laxativa, Pilulæ Cascaræ Compositæ, Pilula Cascara et Belladonnæ et Nucis Vomice, Fluidextractum Rhamni Purshiani Aromaticum, Vinum Rhamni Purshiani.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and US. Not in the others.

Descriptive Notes—The bark of *Rhamnus Purshianus*, DC, as met with in commerce, varies much in appearance and quality. Genuine Cascara Sagrada is more or less externally furrowed longitudinally, and of a reddish-brown tint, with lenticels about $\frac{1}{4}$ inch long, forming slender, whitish, transverse scars in places, it has a characteristic leather-like odour. When the outer surface is scraped the reddish-brown colour of the layer beneath becomes visible, it is of a duller red tint than that of *Rhamnus Frangula*, L. The inner surface is longitudinally striate with projecting medullary

rays, and in some pieces faintly, . . . transversely, and is turned red by alkalis. The *BP* limits the thickness to about $\frac{1}{16}$ inch (1.5 mm), *USP* 1 to 5 mm, which indicates that the thin bark is to be used, probably for the reason that older and thicker bark is usually more bitter. The *USP* directs that the bark should be kept at least one year before being used, but the *BP* does not give any such direction. The recently collected bark is stated to cause vomiting, hence the importance of the mandate. The inner surface varies from pale brown to dark brown, or walnut colour if badly dried. The transverse fracture of the bark is buff coloured or pale brown, but when the bark is kept it becomes darker, that which is thus darkened, as seen in transverse section, is therefore to be preferred, as indicating that it was not recently collected. An inferior variety of thick bark known in the United States as spurious or 'winter' bark, is said to be removed in winter by steaming the branches to soften the bark, and then cutting it off with knives. Another winter form is spoke-shaved, and therefore in chips. The new crop is collected from the end of April till July, and reaches London in August. Sometimes the bark of *R. Californicus*, Esch., and its var. *tomentella*, Benth., is substituted for it, but, according to Rusby, it is only bark that is received from Texas, Arizona, Colorado and New Mexico that is likely to contain it, since the species occurs sparingly only in North California and not in Oregon and Washington, whence supplies of *R. Purshianus* have come during recent years. The chief difference is that the bark of *R. Californicus*, Esch., is of a greyer tint externally, and the lenticels are less numerous and easily become obscured, and the transverse fracture is less dark and more yellow (J. G. Steele) than that of *R. Purshianus*, and the taste is intensely bitter. Under the microscope the bark of *R. Purshianus* is seen to have parallel medullary rays, consisting commonly of two rows of cells, whilst those of *R. Californicus* are shorter, crooked, and not parallel, and are composed of three or more rows (Prescott). The bark of *R. Purshianus* is apt to stain paper yellow, due to the presence of Frangulin. The powdered bark of *R. Purshianus* may be distinguished from that of *R. Frangula* by the presence of sclerenchymatous cells, mucilage is absent. In both, the contents of the parenchymatous cells turn purplish with caustic alkali (Vogl). The bark of Jilhan's *Frangula* has no sclerenchymatous cells and contains mucilage.

Tests—Cascara Bark yields on incineration about 5 p.c. of ash, 8 p.c. should not be exceeded. A limit of 6 p.c. of ash has been suggested. Specimens of good quality bark examined in the author's laboratory showed from 5.6 to 7.8 p.c. of ash.

Preparations

EXTRACTUM CASCARÆ SAGRADÆ. EXTRACT OF CASCARA SAGRADA. *BP Syn*—EXTRACTUM RHAMNI PURSHIANI

Cascara Sagrada, in No. 20 powder, is exhausted by percolation with Distilled Water and evaporated to dryness on a water-bath.

The powdered Bark is moistened with Water and allowed to swell, before it is packed loosely in the percolator

It is an aqueous exhaustion, in *B P* 1885 it was by percolation with Proof Spirit

Dose — 2 to 8 grains = 0 13 to 0 52 gramme

Extractum Rhamni Purshiana (*U S*) is made by exhausting 100 of Cascara Bark in No 60 powder by percolation with Alcohol (12 5 p c), evaporation to dryness, and mixing with sufficient peeled Russian Liquorice Root in No 80 powder to make the product 25

Foreign Pharmacopœias — Official in Belg, Fl, Ital, Mex and U S
Not in the others

EXTRACTUM CASCARÆ SAGRADÆ LIQUIDUM LIQUID
EXTRACT OF CASCARA SAGRADA *B P Syn* — **EXTRACTUM RHAMNI PURSHIANI LIQUIDUM**

5 of Cascara Bark exhausted by percolation with Distilled Water, the percolate evaporated to 3, 1 of Alcohol (90 p c) mixed with 1 of Distilled Water is added, and the whole is made up to 5 by the addition of more Water if necessary

It is almost the same as *B P* 1885

Dose — $\frac{1}{2}$ to 1 fl drm = 1 8 to 3 6 c c

Sometimes given with Ferri et Ammonii Citras and Ammonia

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U S, all with diluted Alcohol U S is 98 p c Alcohol Dan and Swed contain Glycerin Not in the others

Tests — Liquid Extract of Cascara has a specific gravity of 1 055 to 1 070, may contain from 17 to 27 p c w/v of total solids and about 20 p c w/v of Absolute Alcohol

SYRUPUS CASCARÆ AROMATICUS. AROMATIC SYRUP OF CASCARA

Liquid Extract of Cascara Sagrada, 8, Tincture of Orange, 2, Alcohol (90 p c), 1, Cinnamon Water, 3, Syrup, 6
(1 of Liquid Extract in 2 $\frac{1}{2}$)

Dose — $\frac{1}{2}$ to 2 fl dnm = 1 8 to 7 1 c c

Not Official

CAPSULES OF CASCARA — Two strengths, containing concentrated Extract equal to 15 and 30 minims respectively of Fluid Extract

FLUIDEXTRACTUM RHAMNI PURSHIANÆ AROMATICUM (*U S*) — Cascara Sagrada, in No 40 powder, 1000, Glycyrrhiza, in No 30 powder, 100, Magnesium Oxide, 125, macerate with 2000 of Water for 12 hours, then dry it at a gentle heat Percolate this with 250 c c of Glycerin mixed with 500 c c of Alcohol (94 9 p c) and 250 c c of Water, subsequently complete the percolation with diluted Alcohol until exhausted, reserve the first 800 of the percolate and evaporate the remainder to a soft Extract, dissolve this in the reserved portion Add 10 c c of Compound Spirit of Orange, and q s of Diluted Alcohol to make 1000 of fluid Extract

EXTRACTUM CASCARÆ SAGRADÆ LIQUIDUM INSIPIDUM — Liquid Extract of Cascara Sagrada, 100, Liquor Ammoniae Fortior, 7, heat them together on a water bath for 8 hours or until the bitterness has disappeared, and

finally make up the volume to 100 with the requisite quantity of Alcohol (90 p c)
and Water.

The formula given in *BPC* employs 5 of Potassium Hydroxide in place of the Ammonia given above. The *BPC Supplement* gives a formula for a *Miscible Extract of Cascara*. It is the aqueous fluid extract of the *BP* to which Solution of Potassium Hydroxide is added before evaporation, and the Alcohol (90 pc) is replaced by Glycerin. It is stated that the miscible extract should be used in the place of the 'tasteless' liquid extract when making *BPC* preparations.

Dose—30 to 60 minims = 1.8 to 3.6 c c

Many formulas have appeared from time to time, with the object of obtaining a tasteless extract without loss of activity, using either Lime, Magnesia, Potassium Hydride, or Ammonia, we prefer Ammonia, as, apart from the fact that it makes a better preparation, the excess of Ammonia is volatilised, whereas the Potassium Hydride remains in the finished solution. The word 'tasteless' is applied to any of them is a misnomer, and some of the preparations are practically inert. The Bark has a fairly strong flavour peculiar to itself, quite apart from the bitterness.

There is a similar preparation in *Aust.*, named *Extractum Rhamni Purshiani Fluidum*, in which Magnesia is employed.

ELIXIR CASCARÆ *Syn* AROMATIC CASCARA.—Liquid Extract of Cascara Sagrada, 34 5 Liquid Extract of Liquorice, 34 5, Glycerin, 29, Soluble Gluside, 0 75, Oil of Anise, 0 05, Oil of Peppermint, 0 05, Oil of Cloves, 0 025, Oil of Dill, 0 025, Oil of Cinnamon, 0 025, Alcohol, to make 100

This corresponds with the *BPC* formula, it is also given in *Ph. Form* as the Elixir Cascarae Glycerin of the *Australian Ph. Form*

Various formulas for Aromatic Cascara are given in the Hospital Pharmacopœia under the heading *Mistura Cascaræ Composita*, in most of which Liquid Extract of Liquorice forms an important flavouring ingredient.

MISTURA CASCARÆ SAGRADÆ Cascara Mixture.—Liquid Extract of Cascara Sagrada, 1 fl dm, Liquid Extract of Liquorice, 30 minims, Aior " Ammonia, 40 minims, Chloroform Water, to $\frac{1}{2}$ fl oz.—*London*.
 " " of Cascara, 30 minims, Liquid Extract of Liquorice, 30 minims, Aromatic Spirit of Ammonia, 20 minims, Chloroform Water, to 1 fl oz.—*St Thomas's*.

This formula has been incorporated in the *BPC*

MISTURA CASCARÆ SAGRADÆ COMPOSITA
Mixture—Liquid Extract of Cascara, 1 fl. oz., Liquid
30 minims, Sulphate of Soda, 60 grains, Solution of Ammonia, 5 minims,
Water, to 1 fl. oz.—*St. Mary's*

Liquid Extract of Cascara Sagrada, 20 minims, Liquid Extract of Liquorice, 30 minims, Tincture of Belladonna, 5 minims, Tincture of Nux Vomica, 5 minims, Aromatic Spirit of Ammonia, 20 minims, Chloroform Water, *q.s.* to make 1 fl. oz. — *St. Thomas's*

This formula has been incorporated in the *BPC*

Dose — 1 to 1 fl oz = 14 2 to 28 4 c c

Magnesium Sulphate, 60 grains, Glycerin, 1 fl dm, Liquid Extract of Cascara Sagrada, 1 fl drm, Liquid Extract of Liquorice, 60 minims, Tincture of Hyoscyamus, 20 minims, Tincture of Nux Vomica, 5 minims, Compound Decoction of Aloes, to 1 fl oz—*London*

MISTURA CASCARÆ APERIENS—Magnesium Sulphate, 30 grains,
Cascara Sagrada Mixture, to $\frac{1}{2}$ fl oz—*London*

MISTURA LAXATIVA—Liquid Extract of Cascara Sagrada, 1 fl. oz.,
Liquid Extract of Senna, 1 fl. oz., Sodium Bicarbonate, 5 grains, Chloro-
form Water, to 1 fl. oz. In the 1907 edition it is made up to
1 fl. oz. with Water.

PILULA CASCARÆ COMPOSITA (Martindale) — Extract of Cascara, 1½, Extract of Nux Vomica and Alcoholic Extract of Belladonna, of each ½.

Milk Sugar, 1 In grains for one pill, or in grammes for fifteen—*B M J* 93, 11 59b

PILULA CASCARÆ ET BELLADONNÆ ET NUCIS VOMICÆ—Extract of Cascara Sagrada, $\frac{1}{4}$ grain, Extract of Nuc Vomica, $\frac{1}{10}$ grain, Alcoholic Extract of Belladonna, $\frac{1}{10}$ grain Mix and divide into one grain pills—*B P C*

Dose—1 to 3

PILULA CASCARA ET EUONYMIN See EUONYMIN

VINUM RHAMNI PURSHIANI (*Aust.*)—Malaga Wine, 150, Fluid Extract of Cascara Sagrada, 100 Syrup of Orange, 50 Digest eight days and filter Dutch, Cascara Sagrada Bark 1, Malaga Wine 10

CASCARILLA.

CASCARILLA

FR, CASCARILL, GLR, CASCARI LINDB, IIAL, CASCARIGLIA, SPAN, CHACARILLA

The dried Bark of *Croton Eluteria*, J J Bennett

It contains from $\frac{1}{2}$ to 2 p c of an aromatic Oil

Medicinal Properties—Aromatic and stomachic With some physicians it is a favourite bitter tonic Used for the same purposes as Calumba

Prescribing Notes—*The Infusion quickly changes, and will scarcely keep good for a day in summer, but with an aromatic Tincture it keeps well*

The Tincture is frequently prescribed with diluted mineral acids, which, however, usually cause a separation of resin, & fl dim of Mucilage in an 8 oz mixture will keep the resin diffused

Official Preparations—Infusum Cascariellæ and Tinctura Cascariellæ

Not Official—Mistura Cascariellæ Composita, Infusum Cascariellæ Concentratum

Foreign Pharmacopœias—Official in Aust, Dan, Dutch, Ger., Ital, Jap, Norw, Port, Russ, Swed and Swiss Not in Fr, Hung, Mex or Spain An extract is official in Ger

Descriptive Notes—Cascarilla bark varies much in size and quality as found in commerce The best qualities are in quills, or in small curved pieces The outer dull brown or dark grey cork has longitudinal and transverse small cracks and a silver grey surface with minute black dots, a short resinous fracture, and a dark reddish-brown bast, showing thin whitish medullary rays but no sclerenchymatous cells It is bitter and agreeably aromatic The *BP* directions are not quite sufficient to characterise the true bark, since the 'silvery grey patches spotted with minute black dots' occur also in a false bark (referred to *Croton lucidus*, Linn), which causes vomiting and other deleterious effects, and Hartwich has recently shown that of eight Croton barks substituted for Cascarilla since 1901 all agree with it in having 'no groups of sclerenchymatous cells' although differing from Cascarilla bark in other characters (*P J* (4) xxiii p 485) A very slender bark, obtained by speckling the twigs, often occurs in commerce, this is available for pastilles and incense, but is excluded from use in pharmacy by the *BP* description of 'quills from 1 to 3 inches ($2\frac{1}{2}$ to $3\frac{1}{2}$ cm)

or more in length, and from about $\frac{1}{8}$ to $\frac{1}{2}$ inch (4 to 12 mm) in diameter. The general bark is characterised by its aromatic odour and taste, and the greyish-brown layer beneath the whitish coat, where the latter is exfoliated. Under the microscope the powdered bark is distinguished by the characteristic cells containing dark brown secretion, the cork cells thickened chiefly on one side, the absence of stone cells and the presence of starch (Koch). The properties of the bark are due to a bitter principle, resin, and 1.3 p.c. of volatile oil. The ash varies from 6 to 10 p.c. (YBP 1906, p. 209). During recent years it has been deficient in the amount of bitter principle (Naylor, YBP 1906, p. 209). The bark of *Croton lucidus*, which closely resembles that of Cascarilla in appearance, differs in absence of aroma, reddish-brown tint externally, and in the presence of sclerenchymatous cells. Other substitutes differ from true Cascarilla in odour, as well as in the presence of sclerenchymatous cells in the cortex (PJ (4) xv p. 7).

Tests.—Cascarilla leaves about 8 p.c. of ash on incineration, and rarely more than 10 p.c. The average ash of eight samples examined in the author's laboratory showed 8.7 p.c.

Preparations.

INFUSUM CASCARILLÆ. INFUSION OF CASCARILLA

Cascarilla, in No. 10 powder, 1, boiling Distilled Water, 20, infuse for 15 minutes, strain (1 in 20)

Half the strength of BP '85

Dose.— $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 grammes

Incompatibles.—Lime Water and metallic salts

Not in the other Pharmacopœias

TINCTURA CASCARILLÆ. TINCTURE OF CASCARILLA

1 of Cascarilla, in No. 40 powder, percolated with Alcohol (70 p.c.), to yield 5 (1 in 5)

Dose.— $\frac{1}{2}$ to 1 fl. drm. = 1.8 to 3.6 c.c.

Foreign Pharmacopœias.—Aust., Dan., Ital., Jap., Norw., Russ., Swed. and Swiss, 1 in 5. Not in the others.

Tests.—Tincture of Cascarilla has a specific gravity of 0.895 to 0.905, contains from 2 to 3 p.c. w/v of total solids and about 67 p.c. w/v of Absolute Alcohol.

Not Official

MISTURA CASCARILLÆ COMPOSITA.—Tincture of Squill, 10 minims, Compound Tincture of Camphor, 20 minims, Infusion of Cascarilla, to 1 fl. oz.—*Royal Free*

Ammonium Carbonate, 5 grains, Tincture of Squill, 12 minims, Aromatic Syrup, 60 minims, Infusion of Cascarilla, to 1 fl. oz.—*Brompton*

Compound Tincture of Camphor, 15 minims, Vinegar of Squill, 15 minims; Infusion of Cascarilla, to 1 fl. oz.—*St. Thomas's*

This formula has been incorporated in the B.P.C.

INFUSUM CASCARILLÆ CONCENTRATUM.—Cascarilla Bark, in No. 40 powder, 40 parts; Tincture of Cascarilla, 7.5 parts, Alcohol (90 p.c.), 20

parts, Dilute Chloroform Water, 1 in 1000, sufficient to make 100 parts. Prepare by macero expression—*Farr and Wright, P J '06, 1 165 and '07, 1 621, C D '06, 1 252, Y B P 1907, 249*

This formula appears in the *B P C*

Not Official CASSIÆ OLEUM

OIL OF CASSIA

A yellowish or brownish liquid, becoming darker and thicker by age and exposure to the air, having the characteristic odour of Cassia, and a sweetish, spicy, and burning taste. It is a volatile Oil distilled from *Cinnamomum Cassia*.

It should be kept in well stoppered amber tinted glass bottles, in a cool place, and away from the light.

The principal constituent of Cassia Oil is Cinnamic Aldehyde, of which it should contain at least 75 p c. Cinnamyl Acetate and traces of Cinnamic Acid are also present. A stearoptene, Ortho methyl-coumaric aldehyde, has been shown to be a constituent of the oil.

Soluble in an equal volume of Alcohol, the solution being slightly acid to Litmus paper, also soluble in an equal volume of Glacial Acetic Acid.

This Oil is official in the German and U S Pharmacopœias under the name 'Oleum Cinnamomi'.

Medicinal Properties—It possesses the aromatic, carminative and anti septic properties of Cinnamon bark, but the oil is a powerful local stimulant.

Dose— $\frac{1}{2}$ to 3 minims = 0.08 to 0.18 c c

Foreign Pharmacopœias—Official in Belg, Ger, Jap, Norw, Swiss and U S

Tests—Cassia Oil has a specific gravity of 1.053 to 1.065. It should be optically almost inactive, and should never vary more than one degree to the right or left in a tube of 100 mm. It should be soluble in 3 to 4 parts of Alcohol (70 p c), and in all proportions in Alcohol (90 p c).

When shaken with an equal volume of Nitric Acid at 0° C (32° F), a crystalline mass should result. With Ferric Chloride Solution the alcoholic solution of the oil should give a brown coloration. The oil forms a crystalline compound with Sodium Bisulphite Solution (30 p c), and this reaction is utilized as a means of determining the percentage of Cinnamic Aldehyde. A measured quantity of 10 c c of the oil is shaken in a flask with a long thin neck graduated to one-tenth c c, with 10 c c of a 30 p c Sodium Bisulphite Solution, and warmed in a water bath until the contents are liquefied. When this has occurred more Sodium Bisulphite Solution is added, the mixture being constantly heated and occasionally shaken until the flask is quite three-fourths filled. The heating in the water-bath is continued until no solid particles are visible, and the odour of Cinnamic Aldehyde has disappeared. When this is effected the contents of the flask are allowed to cool, and sufficient of the 30 p c Sodium Bisulphite Solution added to bring the lower level of the oily layer to the zero mark on the graduated neck of the flask, and the number of c c is read off. The *U S P* requires that the residual liquid should not measure more than 2.5 c c, indicating at least 75 p c by volume of Cinnamic Aldehyde. The *P G* process is to take a measured quantity of 5 c c of the oil, and after mixing it with 45 c c of a 30 p c Sodium Bisulphite Solution, to heat it in a water bath for two hours with intervals of frequent shaking. Not more than 1.5 c c of oil shall remain undissolved, indicating at least 70 p c by volume of Cinnamic Aldehyde.

The more generally occurring sophistications are Colophony Resin and Petroleum, Lead and Copper. For the detection of Colophony Resin and Petroleum, the *U S P* mixes 1 c c of the oil with 8 c c of a mixture of 3 volumes of Alcohol (94.9 p c) and 1 volume of Water, when a clear solution should result, and if to this solution 2 c c of a saturated Lead Acetate Solution in a mixture of 3 volumes of Alcohol (94.9 p c) and 1 volume of Water be added,

no precipitate should be produced. The *PG* includes a somewhat similar test, dissolving the oil in 3 to 4 parts of Alcohol (90 p c), and treating the solution with half its volume of freshly-prepared Lead Acetate Solution. An additional test for Resin is given in the latter Pharmacopœia, 100 parts of the oil, heated on a water-bath until all volatile constituents have been dissipated, should not yield more than 8 parts of residue. The value of this test may be much enhanced by performing it in a tared fractionating flask, and recording the temperature at which the distillate passes over. A quantity of 50 grammes of the oil should be weighed into the flask, and the oil distilled over a direct flame. After the Water has passed over, the thermometer rapidly rises to 240°C (464°F), and the major portion of the oil passes over between 240°C (464°F) and 260°C (500°F). The appearance of white fumes indicates the end of the distillation, the residue should be viscid and tough, and not hard and brittle. The presence of Petroleum is detected by the solubility of the distillate in Alcohol (70 p c). The presence of Lead and Copper is due to the solvent action of the Cinnamic Acid produced by the Aldehyde oxidation on the metal of the containing vessels, and is readily detected by Hydrogen Sulphide. Lead is chiefly present in unrefined oils. The *PG* includes a colour test with Ferric Chloride Solution for Phenol, requiring that the colour produced on the addition of the Ferric Chloride Solution shall be brown, and not a green or blue.

CINNAMIC ALDEHYDE ($\text{C}_9\text{H}_8\text{O}$, eq 131.07) — This Aldehyde is the principal constituent of oils of Cassia and Cinnamon. It may also be prepared artificially by the action of Sodium Hydroxide upon a mixture of Benzaldehyde and Acetic Aldehyde.

A clear, colourless, or pale yellow, highly refractive liquid possessing a characteristic aromatic odour and a sweetish spicy and subsequently burning taste. It should contain not less than 95 p c of pure Cinnamic Aldehyde.

Its use in medicine is similar to that of Oil of Cassia.

Dose — $\frac{1}{2}$ to 2 minims = 0.03 to 0.12 c c.

Foreign Pharmacopœias — Official in Austria, Sweden and U.S.

Tests — Cinnamic Aldehyde has a specific gravity of 1.054 to 1.056, and boils about 247°C (476.6°F). It is optically inactive. It should conform to the tests given under Cassia Oil. It forms a crystalline compound with Sodium Bisulphite, and when mixed with an excess of a 30 p c solution of the salt should dissolve completely leaving practically no oily residue.

It should be free from the sophistications mentioned under Oils of Cassia and Cinnamon.

The *USP* method of determination is to introduce 12 drops of the Aldehyde into a carefully counterpoised 150 c c flask and carefully ascertain the exact weight. 5 c c of Distilled Water and a few drops of Rosolic Acid Solution are added and the solution exactly neutralised by the addition of Tenth-normal Volumetric Sodium Hydroxide Solution. A measured quantity of 50 c c of a 20 p c Sodium Bisulphite solution is then added, and the flask is immersed in a water-bath of boiling Water. Sufficient Semi-normal Volumetric Hydrochloric Acid Solution is added to maintain the neutrality of the liquid and a drop or two of Rosolic Acid Solution, the flask being kept continuously heated and frequently agitated. The number of c c of Semi-normal Hydrochloric Acid Solution used is noted when a permanent state of neutrality is reached. A blank test with the same materials without the Cinnamic Aldehyde is used as a control, and the number of c c of Semi-normal Volumetric Hydrochloric Acid used in the blank is subtracted from the number of c c used in the original. Each c c of the difference corresponds to 0.083 gramme Cinnamic Aldehyde.

Not Official

CASSIA BEAREANA

A small tree, attaining the height of 20 or 30 feet growing in equatorial East Africa. A decoction of the roots has been recommended in the treatment of

blackwater fever and in hæmaturia. The decoction is prepared by the natives by boiling about a dozen pieces of the root, about 1 inch long, in a gallon of Water, and it is administered in teacupful doses. The powdered bark is applied as a dressing to ulcers.—*L* '02, i 283, '03, i 190, *P J* '01, ii 616, '02, i 42 *C D* '03, i 872

A fluid extract (1 in 1) is also made, dose 30 to 60 minims = 1.8 to 3.6 c c

CASSIÆ PULPA.

CASSIA PULP

FR, PULPE DE CASSI, GER, ROHRENKASSIE, ITAL, CASSIA, SPAN,
CANAFISTULA

The Pulp from the Fruits of *Cassia Fistula*

Imported from the East or West Indies

Medicinal Properties—Laxative. Useful in small doses for habitual constipation. Large doses occasion nausea, flatulence, and griping, generally given in combination, as in Confection of Senna.

Dose—60 to 120 grains = 4 to 8 grammes, as a laxative, 1 to 2 oz = 28.4 to 56.8 grammes, as a purgative.

Official Preparation—Contained in Confectio Sennæ, 1 part in 8 nearly.

Foreign Pharmacopœias—Austri, Fruit and Pulp, Belg, Fruit, Ital, Mex, Port and U S, Fruit. Not in the others.

Descriptive Notes—The fruit of the *Cassia Fistula*, L, is a cylindrical, indehiscent pod, separated by thin internal transverse partitions into numerous cells each of which contain a single seed, immersed in a blackish pulp. The pods are chiefly imported from Dominica in the West Indies, but those from the East Indies, which are smaller, smoother, and have a blacker pulp are usually preferred, the pulp of this kind being considered more active. Some of the East Indian pods come from Sourabaya in Java, *vid* Amsterdam. Pods in which the seeds rattle when shaken are considered old and inferior. The pulp usually forms about 30 p c of the weight of the pods. The official description, *viz*, '1½ to 2 ft long (35 to 50 cm) and from ¼ to 1 in (18 to 25 mm) in diameter, the sutures being marked by two smooth longitudinal bands,' excludes the fruits of *Cassia grandis*, L, which has larger compressed pods, of which the ventral suture is marked by two prominent marginal ridges, and the surface of the pods has prominent veins. It also excludes the smaller fruits of *Cassia moschata*, H B and K, which have a brown pulp. The term viscid applied to the pulp implies that old pods with hard dry pulp should not be used. The extracted pulp will not keep in the viscid condition, soon becoming mouldy, especially in a damp place, and is therefore sometimes met with in commerce in the form of tough extract not easily miscible with the other ingredients of Confectio Sennæ unless first rubbed down with Water. It should therefore be prepared fresh for this purpose. Cassia pulp is said to be one of the ingredients used in Turkey for adulterating Opium.

Not Official

CASTOREUM

FR, CASTOREUM, GER, BIBERGEIL, ITAL, CASTOREO, SPAN, CASTORLO

The dried preputial follicles and then secretion, obtained from the Beaver, *Castor Fiber*, L, the oil sacs being rejected

Medicinal Properties—Moderately stimulant and antispasmodic, occasionally used in hysteria and spasmodic disorders

Dose—Of the powder 5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes—*The Tincture when mixed with Water will yield a deposit after a time, it should therefore be prescribed with Mucilage of Gum Acacia, 8 fl drms in 8-oz mixture*

Foreign Pharmacopœias—Official in all except Belg, Dan, Dutch, Ger, Jap, Swed and U.S. Both the Canadian and Russian varieties are official in Russ. Austri not more than 40 p.c. insoluble in hot Alcohol, and not more than 4 p.c. of ash

Descriptive Notes—The Castoreum of commerce is chiefly imported from the Hudson's Bay Territory. It consists of two pyriform sacs about 2 in (5 cm.) long, usually compressed and wrinkled, and one of the sacs is rather longer than the other. The contents of the sac are of a reddish-brown colour, resinous, and softening readily when warmed. Castoreum has a characteristic odour. It is liable to deteriorate unless kept quite dry.

TINCTURA CASTOREI—Castor, in coarse powder, 1, Alcohol (90 p.c.), 20, macerate seven days, agitating occasionally, strain, press, and add Alcohol *q.s.* to yield 20 (1 in 20)

Dose— $\frac{1}{2}$ to 1 fl. drim = 1.8 to 3.6 c.c.

Foreign Pharmacopœias—Official in Austri, Hung, Norw and Port, 1 in 5, Fr, Ital, Russ and Swiss, 1 in 10, Mex, 1 and 10, Span, 1 in 25, all by weight. Not in Belg, Dan, Dutch, Ger, Jap, Swed or U.S.

Russ contains a Tincture made with Russian Castor, and also one made with Canadian Castor

CATECHU.

CATECHU

B.P. Syn—CATECHU PALLIDUM

FR, CACHOU, GER, KATECHU, ITAL, CATECU, SPAN, CATECÚ

An extract of Leaves and young Shoots of *Uncaria Gambier*, Roxb.

It contains from 80 to 40 p.c. of Catechu-Tannic Acid, from 10 to 40 of Catechin, some mucilage and mineral matter

Prepared in Singapore and in other places in the Eastern Archipelago

Terra Japonica is a trade term (now almost obsolete) applied both to Cutch and Gambier

Solubility.—Almost entirely soluble in boiling Water. 70 to 75 p.c. is soluble in Alcohol (90 p.c.) 50 to 60 p.c. is soluble in cold Water, and the solution is bright

Medicinal Properties.—A powerful astringent. Used in diarrhœa, dysentery, gastric and intestinal hæmorrhage and for other purposes for which Tannic Acid is used. Lozenges are the best medium for administering it in relaxed conditions of the throat and in ulcers of the mouth

Dose.—5 to 15 grains = 0.32 to 1 gramme

Incompatibles—The Alkalis, metallic salts, and Gelatin.

Official Preparations—*Pulvis Catechu Compositus*, *Tinctura Catechu*, and *Tiochiscus Catechu*

Not Official—*Mistura Catechu et Cretæ*, *Mistura Hæmatoxyli cum Catechu*

Foreign Pharmacopœias—Official in Dutch, Ger., Jap. and Port (Cato)
Not in the others See below, *CATECHU NIGRUM*

Descriptive Notes—Under the commercial name of Gambier, or more rarely *Terra Japonica*, several qualities of the drug may be met with. The inferior kinds are largely used for tanning, dyeing and calico printing. The form official in the *B.P.* consists of cubes about 1 in (25 mm) in diameter, reddish brown externally, and pale cinnamon-brown and porous and friable internally. Smaller cubes are also met with in commerce and occasionally parallelograms and discs, or lozenges with fluted margins, but the last two are usually paler in colour and contain Starch. They are used in India for chewing with the Betel pepper leaf, but are not admissible for pharmaceutical use, since official *Catechu* should not afford any characteristic reaction with the tests for Starch.

Tests—*Catechu* is required officially to be almost completely soluble in boiling Water, and to be soluble to the extent of 70 p.c. in Alcohol (90 p.c.)

It should not yield a pronounced blue coloration on the addition of Iodine Solution to its boiled and cooled aqueous solution, and it is officially required to yield not more than 5 p.c. of Ash when incinerated. A sample examined in the author's laboratory left 4.0 p.c. of ash. The presence of Black *Catechu* may be detected by the marked green fluorescence produced in the Petroleum Ether solution, when 3 grammes of *Catechu* are mixed with 25 c.c. of Normal Volumetric Sodium Hydroxide Solution and shaken with 50 c.c. of Petroleum Ether.

The extract official in the *U.S.P.* is prepared from *Ououparia Gambu*, Baillon, that of the *P.G.* from either *Ououparia Gambu* or *Acacia Catechu*, Willd. The *P.G.* states the highly-diluted alcoholic solution gives a green coloration with Ferric Chloride T.S. The residue insoluble in Water, after washing with hot Water, should when dried at 100° C (212° F) not amount to more than 15 p.c. The *P.G.* ash limit is 6 p.c., that of the *U.S.P.* not more than 5 p.c.

Preparations

PULVIS CATECHU COMPOSITUS COMPOUND POWDER OF
CATECHU

Catechu, 4, Kino, 2, *Klameia* Root, 2, Cinnamon Bark, 1,
Nutmeg, 1 (1 in 2½)

Keep it in a stoppered bottle

Dose—10 to 40 grains = 0.65 to 2.6 grammes

Not in the other Pharmacopœias, a powdered Black *Catechu* is official in Spain

TINCTURA CATECHU TINCTURE OF *CATECHU*

Catechu, in coarse powder, 4, Cinnamon Bark, bruised, 1,
Alcohol (60 p.c.), 20, prepare by the maceration process

Now 1 in 5 instead of 1 in 8

Dose — $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 cc

Foreign Pharmacopœias —Official in US (Tinct Gambii Co), 1 in 20, Belg, Fl (Teinture de Cachou), Dutch, Ger, Ital, Jap, Port and Swiss, 1 in 5, Mex, 1 and 5, all by weight (except US), and with the exception of the Dutch, with **Black Catechu**. Not in the others.

Tests —Tincture of Catechu has a specific gravity of 0.978 to 0.983, should contain 15.5 to 18 p.c. w/v of total solids and about 52 p.c. of Absolute Alcohol. It yields a dirty green coloration with TS of Ferric Chloride.

TROCHISCUS CATECHU. CATECHU LOZENGE

1 grain of Catechu in each, with Simple Basis

Dose —1 to 6 lozenges

Foreign Pharmacopœias —Official in US, 1 grain Gambii in each, Fr and Dutch, about $1\frac{1}{2}$ grains in each. Not in the others.

Not Official

MISTURA CATECHU ET CRETÆ —Tincture of Catechu, 20 minims, Chalk Mixture, to 1 fl oz —*University*
Tincture of Catechu, 4, Chalk Mixture, q.s. to produce 100 —*B.P.C. Supplement*

MISTURA HÆMATOXYLI CUM CATECHU —Tincture of Catechu, 40 minims, Diluted Sulphuric Acid, 15 minims, Decoction of Logwood, to 1 fl oz —*St. Thomas's*

This formula has been incorporated in the *B.P.C.*, using Aromatic Sulphuric Acid.

Not Official

CATECHU NIGRUM

BLACK CATECHU

A dried extract from the wood of *Acacia Catechu*, Willd. It is also known as **Pegu Catechu** and **Cutch**. It generally occurs in irregularly shaped blackish-brown masses, astringent, and bitter in taste. The *Ind* and *Col Add* put the limit of ash at 6 p.c.

Solubility —About 80 to 90 p.c. is dissolved by cold Water, the solution being very turbid. The *Ind* and *Col Add* requires not less than 80 p.c. to be soluble in Alcohol (90 p.c.)

Dose —5 to 15 grains = 0.32 to 1 gramme

Official in the *Ind* and *Col Add* for India, Eastern Colonies and North American Colonies, within which it may be used in making the *B.P.* preparations for which Catechu is ordered.

Foreign Pharmacopœias —Official in Austri, Belg, Fl (Cachou de Pégu), Ger, Ital, Jap, Mex, Port (Cato), Russ, Span and Swiss, US **Ourouparia Gambir**. Not in the others.

Tests —The aqueous solution is acid in reaction towards blue Litmus paper, and yields a green coloration with Ferric Chloride Test-solution, the colour changing to purple on the addition of Sodium Hydroxide Solution. About 80 to 90 p.c. should be dissolved by cold Water, and in boiling Water it should be almost completely soluble. The matter insoluble in Alcohol (90 p.c.) should not amount to more than 20 p.c. The *P.G.* states that the residue insoluble in Water, after washing with hot Water, shall when dried at 100° C (212° F) not amount to more than 15 p.c. The ash should not amount to more than 6 p.c. Two samples examined in the author's laboratory left 4.2 p.c. and 5.0 p.c. of ash.

Not Official CAULOPHYLLUM

BLUE COHOSH

The Rhizome and Roots of *Caulophyllum thalictroides*, Mich

Descriptive Notes —The rhizome is of a greyish brown colour, knotty, $\frac{3}{4}$ to 4 inches (7.5 to 10 cm) long and $\frac{1}{4}$ to $\frac{1}{2}$ inch (6 to 8 mm) in diameter, with few branches, and broad concave stem scars at short intervals, the internodes being marked with transverse rings about $\frac{1}{16}$ to $\frac{1}{8}$ inch (1.5 to 3 mm) apart, and is furnished with matted undulated rootlets, about 4 inches (10 cm) long and 1 mm broad. The transverse section is whitish and often porous. The bark is thin, the woody wedges short, the medullary rays broad, and the pith large. The rootlets have a relatively thicker bark, and a tough woody centre. It contains starch. It is almost free from odour, and has a sweetish bitter and somewhat acid taste. The distinctive microscopic characters are the small, spherical, simple starch granules, the large and small porous vessels and the tracheids.

Caulophyllum contains a body analogous to Saponin, and termed for distinction Leontin, and a colourless, odourless, and tasteless, non-crystalline alkaloid, Caulophylline, which is soluble in Water, Alcohol (90 p c), Ether and Chloroform, and which on treatment with Hydrochloric Acid is converted into a crystalline salt, **Caulophylline Hydrochloride**.

Caulophyllin, an eclectic remedy in the form of a brown powder, has been recommended as an emmenagogue, sedative and diuretic. Also employed with success as an anthelmintic. Given in the form of a 1 in 20 decoction or infusion, or as a 1 in 5 tincture. Best given in form of a pill.

Dose —1 to 4 grains = 0.06 to 0.26 gramme. Of the decoction or infusion, 1 to 2 fl oz = 28.4 to 56.8 c c. Of the tincture, 1 to 2 fl drms = 3.6 to 7.2 c c.

A **fluid extract** (1 in 1) and a **compound fluid extract** are also given in doses of 30 to 60 minims.

FLUIDEXTRACTUM CAULOPHYLLI —1000 grammes of Caulophyllum, in No. 60 powder, exhausted by percolation with Alcohol (70 p c) to produce 1000 c c of fluid extract.

This was incorporated in the *B P C* employing Alcohol (60 p c), but was changed in the *B P C Supplement* to Alcohol (70 p c).

LIQUOR CAULOPHYLLI ET PULSATILLÆ (*Ph Form*) —Caulophyllum root (Blue Cohosh), 10 oz, Pulsatilla, 10 oz, Rectified Spirit, a sufficiency, Water, a sufficiency. Macerate the coarsely ground drugs in 60 oz of Rectified Spirit for forty-eight hours and transfer to a percolator. Reserve the first 12 oz of percolate and continue percolation with 60 oz of Water. Recover the spirit from this percolate and evaporate to 8 oz. Mix this with the reserved portion, acidify with dilute Sulphuric Acid 80 minims ($\frac{1}{8}$ fl oz), set aside for a day, and filter.

This formula has been incorporated in the *B P C*, but the formula has been altered in the *B P C Supplement* to Liquid Extract of Caulophyllum, 25, Liquid Extract of Pulsatilla, 5, Glycerin, 15, Alcohol (60 p c), *qs* to make 100.

A formula is also inserted in the *Supplement* for Liquor Caulophylli et Pulsatillæ Compositus.

CERA FLAVA

YELLOW BEESWAX

FR, CIRE JAUNE, GER, GELBES WACHS, ITAL, CERA GIALLA, SPAN, CERA AMARILLA

A hard, yellow, or yellowish-brown, waxy solid, formed by the Hive-Bee, *Apis mellifica*, L.

When quite fresh, is of a golden yellow, but on keeping gets darker.

Solubility.—Entirely in Oil of Turpentine insoluble in Alcohol (90 p c), slightly, and not uniformly, soluble in (cold) Ether, about 1 in 100 of boiling Alcohol (90 p c), 1 in 10 of boiling Ether

Medicinal Properties.—Chiefly used in medicine as an ingredient of plasters and ointments, and is preferable to white Beeswax as the ointments made with the yellow keep a longer time without becoming rancid

Official Preparation—Cera Alba. Used in the preparation of Emplastrum Calafaciens, Emplastrum Cantharidis, Unguentum Menthol, Emplastrum Picis, Unguentum Hydragyri Compositum, Unguentum Picis Liquidæ, Unguentum Resinæ and Unguentum Staphisagriæ

Not Official—Aseptic Wax

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex (Cera Amarilla), Norw., Port (Cera Amarilla), Russ., Span., Swed., Swiss and U S

Descriptive Notes—Yellow Beeswax is imported from many parts of the world, and as it is not in all cases produced by *Apis mellifica*, but by other species in Africa and India, it might become a local question whether any Beeswax except that of *Apis mellifica* is intended by the Pharmacopœial description. This difficulty might have been avoided by the addition of the words 'of other species'. The wax imported from East Africa often shows a minutely crystalline surface which appears to be characteristic of it

The value of Beeswax depends on its purity. Owing to the use of artificial foundations for the honeycomb, a certain percentage of paraffin wax, Paraffin is often found in Beeswax as an impurity. Some Beeswax is very impure and dark in colour, due to imperfect straining, and it is also sometimes deficient in odour

Tests.—The distinguishing tests for Beeswax are that it has a specific gravity of about 0.960, the *BP* gives 0.960 to 0.970, the *USP* 0.951 to 0.960 at 25°C (77°F), the *PG* 0.962 to 0.966, the method adopted by the latter Pharmacopœia for ascertaining the specific gravity being given in the small type below, a melting point of 61° to 64°C (141.8° to 147.2°F), the *USP* gives 62° to 64°C (143.6° to 147.2°F), the *PG* 63° to 64°C (145.4° to 147.2°F), the method adopted by the *BP* for the determination of the melting point being given in the small type below under that heading, and a solidifying point of 58° to 61°C (136.4° to 141.8°F). It is officially required to possess an Acid value of not less than 17.8, an Ester value of from 69.08 to 75.76, and a Saponification value of from 86.88 to 93.56. The *USP* gives no Acid or Ester value, but requires a Saponification value of from 90 to 96. The *PG* gives an Acid value of from 18.5 to 24.15, an Ester value of from 73.0 to 75.8, and a Saponification value of from 94.3 to 97.15. The inclusion of an Iodine absorption figure would have been an advantage, genuine Beeswax absorbing from 8 to 9 p c of Iodine. The methods adopted by the various Pharmacopœias for the determination of the Acid, Ester, and Saponification values are given below in small type. Samples of laboratory showed

Acid values ranging from 20.0 to 22, with an average of 21.2, Ester values from 68 to 77, with an average of 72.6, Saponification values of 88 to 99, with an average of 91.3, and a fairly constant Iodine value of 8.9 p.c.

The more generally occurring adulterants are Paraffin or Ceresin Wax, Fatty Acids and Tallow, Japan Wax and Resin, Soap, Starch, and insoluble matter. The solubility of the Wax in Water detects the presence of Soap, and the solubility in boiling Sodium Hydroxide Solution detects acids, *e.g.*, Stearic Acid, Tallow, Japan Wax, and Resin. The resulting solutions should neither be turbid nor when acidified with Hydrochloric Acid should they yield precipitates. Paraffin and Ceresin Wax are detected by heating the Wax with 5 times its weight of Sulphuric Acid for 15 minutes and diluting the mixture with Water. The *BP* says that no solid, wax-like body should separate, the *USP* says no notable amount of solid substance 'which cannot be decomposed by Sulphuric Acid on further treatment' should separate. The solubility of the Wax in Oil of Turpentine detects the presence of Starch and mineral matter. The washed residue when treated with boiling Water, cooled, and tested with Iodine solution shall yield no characteristic blue coloration. Stearic Acid and Resin increase the Acid value and decrease the Ester value, Paraffin and Ceresin Wax decrease both the Acid and Ester value, Carnauba Wax decreases the Acid value but has no effect upon the Ester value, Japan Wax has no influence upon the Acid value but increases the Ester value, whilst Tallow and vegetable Wax increase the Ester value. Stearic Acid, Paraffin and Ceresin Wax lower the Iodine absorption, whilst Resin and Tallow increase it.

Specific Gravity—The *PG* gives the following directions for taking the specific gravity. Let 2 parts of Alcohol be mixed with 7 parts of Water and the mixture allowed to stand at 15° C (59° F), until all air bubbles have disappeared from it, then let a small ball of yellow wax be introduced into it. This wax should float on the liquid or at least be suspended when the specific gravity of the diluted Alcohol has been brought to from 0.962 to 0.966 by the addition of Water. The ball of Beeswax requisite for this should be prepared as follows. Let the wax be melted at as low a temperature as possible and allowed to fall drop by drop into a beaker containing Alcohol. Before the fully-rounded mass so obtained is used for the determination of the specific gravity it should remain in the air for twenty-four hours, *PG*.

Melting Point—The *BP* method is to draw up into a capillary tube of 1 mm internal diameter some of the melted wax, and allow it to cool for three hours before taking the melting point. The tube is fitted to the bulb of a thermometer and both are immersed in Water contained in a glass vessel, which must be heated gradually until the wax liquefies.

Sodium Hydroxide—The *USP* gives the following test. If 1 gramme of yellow wax be boiled for half an hour with 35 c.c. of an aqueous solution of Sodium Hydroxide (1 in 7), the volume being preserved by the occasional addition of Water, the wax should separate on cooling without rendering the liquid opaque, and no precipitate should be produced in the liquid after filtration through glass wool or asbestos on the addition of Hydrochloric Acid (absence of fats, fatty acids, Japan Wax, and Resin). This test is also given in the *BP*, but no quantities are stated. In the *PG* test 1 gramme of Yellow Wax is boiled with 10 c.c. of Water and 3 grammes of Sodium Carbonate. The Wax should separate out over the saline solution on cooling, and the solution should be rendered not more than opalescent.

Hydrochloric Acid—Hydrochloric Acid should not produce a precipitate in Water which has been boiled with a portion of Yellow Wax, *U S P* and *B P*, but the latter does not specify boiling Water

Litmus—If 1 gramme of Yellow Wax be boiled for a few minutes with 20 c c of Alcohol and filtered after an hour, the cooled almost colourless liquid should neither redden blue Litmus paper nor after adding Water to it should it become strongly turbid, *P G*

Acid Value—5 grammes of Beeswax require for neutralisation of the acids not less than 16 c c of Normal Volumetric Alcoholic Solution of Potassium Hydroxide, using Phenolphthalein Solution as an indicator, *B P*, the *P G* direct the use of 5 grammes of the Beeswax, 50 c c of Alcohol, and after the addition of 20 drops of Phenolphthalein Solution, titration with Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution, of which from 3.3 to 4.3 c c should be necessary for neutralisation

Ester Value—To the residual mixture left after the *B P* determination of the Acid value, a measured quantity of 20 c c of Normal Volumetric Alcoholic Potassium Hydroxide Solution is added, and the mixture saponified under a reflux condenser for one hour, the excess being titrated with Normal Volumetric Sulphuric Acid Solution, for which from 13.2 to 13.8 c c should be necessary, indicating that not less than 6.2 nor more than 6.8 c c has been absorbed by the Beeswax. In the *P G* test, after the further addition of 20 c c of Semi-normal Potassium Hydroxide Solution to the residue from the Acid value determination the mixture is warmed for half an hour on a water-bath, and the excess alkali is then titrated with Semi-normal Volumetric Solution of Hydrochloric Acid, when from 6.5 to 7 c c of the acid should be necessary for the neutralisation of the uncombined alkali

Sulphuric Acid.—If 5 grammes of Beeswax are heated with 25 c c of Sulphuric Acid for 15 minutes to 160° C (320° F), and the mixture diluted with Water, no solid wax-like body should separate, *U S P*, *B P* uses 25 grammes

Saponification Value—Yellow Wax saponified by Alcoholic Potassium Hydroxide T S should show a Saponification value of 90 to 96, these figures representing the number of milligrammes of Potassium Hydroxide required to saponify one gramme of Wax, *U S P*

Not Official

ASEPTIC WAX—Beeswax, 87, Almond Oil, 12, Salicylic Acid, 1. Melt the Beeswax and Oil, strain through muslin, add the Salicylic Acid, heat to 150° C (300° F) in an oil bath, allow to cool slightly, pour into stoppered bottles, which have been sterilised, and when cold add to each bottle sufficient aqueous solution of Mercuric Chloride (1 in 500) to cover the Beeswax

This Wax is made by us for Sir Victor Horsley, who uses it for arresting hæmorrhage from cranial bones, by smearing it over the bleeding surface. Put up in wide-mouthed stoppered bottles under aseptic conditions

This formula has been incorporated in the *B P C*

CERA ALBA.

WHITE BEESWAX

FR, CIRE BLANCHE, GER, WEISSES WACHS, ITAL, CERA BIANCA, SPAN, CERA BLANCA

A white, or almost white, waxy, somewhat translucent solid, or in thin circular white cakes, obtained by melting yellow Beeswax

Official Preparations.—Contained in *Pilula Phosphori*, *Suppositoria Acidi Carbolic*, *Unguentum Aquæ Rosæ*, and *Unguentum Cereæ*

Not Official.—*Ceratum*, *Unguentum Simplex*.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Norw, Port (Cera Branca), Mex, Russ, Span, Swed, Swiss and U S

Descriptive Notes—The White Beeswax of the *B P* should consist of English, or at least of European Beeswax, since it is Yellow Beeswax made by the *Apis mellifica*, Linn, which has been bleached by exposure to moisture, air and light. The Beeswax sold in commerce in blocks under the name of Dentists' Wax is the purest obtainable, but usually has a faint yellowish tinge. That in flat circular cakes commonly contains 1 to 2 p c of Paraffin Wax, which is probably due to the fact that foundations made with Paraffin are laid down in hives for the bees to build upon.

Tests—Officially it is required to correspond to the tests for Yellow Beeswax, but during the process of bleaching by chemical means changes occur in the composition of the Wax which cause an alteration in the physical constants. It has a specific gravity of 0.960 to 0.970, a melting point of 62° to 64° C (143.6° to 147.2° F), an Acid value from 19 to 23, an Ester value from 74 to 84, a Saponification value from 93 to 107, and an Iodine absorption 1 to 10 p c.

Samples examined in the author's laboratory gave Acid values ranging from 11.8 to 24.6, with an average of 15.7, Ester values ranging from 71.7 to 100.8, with an average of 92, Saponification values ranging from 96 to 114.8, with an average of 107.7, the Iodine values ranged from 2.5 to 7.6, with an average of 5.5.

The more generally occurring adulterants are those mentioned under *Cera Flava*, and the same methods as there described may be employed for their detection.

Not Official

CERATUM (U S)—White Wax, 6, White Petrolatum, 4, Benzoinated Lard, 10, melt until the mixture is liquefied, and stir it constantly until it congeals. For use in southern latitudes, and during the heated season in other localities 1 of Benzoinated Lard may be replaced by an equal quantity of White Wax.

UNGUENTUM SIMPLEX—Formerly Official in *B P*, but now omitted.

Foreign Pharmacopœias—Official in Austr and Hung, Lard 8, White Wax 2, Belg, Lanolin 5, Vaseline 5, Dutch, yellow Wax 3, Sesame Oil 7, Mex, White Wax 1, Sesame Oil 3, Swiss, White Wax 30, Olive Oil 70, Ethereal Tincture of Benzoin 10, U S Benzoinated Lard 8, Yellow Wax 2, Ger (Unguentum Cereum), Olive Oil 7, Yellow Wax 3, Jap, Yellow Wax 1, Sesame Oil 2, Port (Cerato Simplex), White Wax 3, Almond Oil 7, Span (Cerato Simplex), White Wax 1, Almond Oil 3, Swed (Ceratum), White Wax 1, Spermaceti 1, Benzoinated Lard 3, Norw (Ung Cereum), and Russ (Ung Cereum), Olive Oil 3, Yellow Wax 1.

CERII OXALAS.

CERIUM OXALATE

$\text{Ce}_2(\text{C}_2\text{O}_4)_3, 10\text{H}_2\text{O}$, eq 719.22

A white, or almost white, odourless and tasteless granular powder. Officially it is stated to be prepared by the decomposition

of a soluble salt of Cerium with a soluble Oxalate. It has been prepared from the minerals Cerite and Thorite, but is now more generally obtained from Monazite. It consists principally of Cerium Oxalate. Commercial samples usually contain the Oxalates of Lanthanum and Didymium and of other rare earths of this group.

It has been pointed out (*C D* '00, 1 992) that the formula should show 10 molecules of Water of crystallisation, and not 9 as in the official formula, and that pure Cerium Oxalate free from Didymium yields a pure yellow, and not a brownish-red residue on ignition.

In spite of two references to this point in *Pharmacopœia Committee Reports* it is interesting to note that the *BP C* still retains the $9H_2O$ formula.

Solubility—Insoluble in Water, Alcohol (90 p c) and in Ether. Insoluble in cold, but decomposed by boiling Potassium or Sodium Hydroxide Solutions. Insoluble in cold, but soluble in hot, diluted Sulphuric or Hydrochloric Acid.

Medicinal Properties—Gastric sedative. Given in chronic vomiting, and vomiting during influenza, and of phthisis, also in dyspepsia, gastrodynia, and pyrosis. It has been recommended in sea-sickness, in doses of 10 to 20 grains every three hours. Given with success in spasmodic cough of gastric origin.

Cerium Oxalate in the gastric crisis of *Tabes*—*L* '96, 11 551.

Dose.—2 to 10 grains = 0.13 to 0.65 gramme.

Prescribing Notes—*It is taken in 5 to 15 grain doses as a powder mixed with a little Water, also given in cachets. It is also supplied in effervescent granules containing 1, 2, 3 and 5 grains in each diachm.*

Foreign Pharmacopœias—Official in Jap, Mex and Port (Oxalato de Cerio), U S. Not in the others.

Tests—The distinguishing tests for Cerium Oxalate are that at a dull red heat it is decomposed, leaving a reddish-brown powder, which is completely soluble in boiling Hydrochloric Acid, the solution yielding on the addition of a saturated Potassium Sulphate solution a white crystalline precipitate, Potassium or Sodium Hydroxide Solution yields a white precipitate which gradually turns yellow on contact with the air, and which does not dissolve in an excess of the reagent, whilst Ammonium Carbonate Solution yields a white precipitate more or less soluble in an excess of the reagent. In carrying out the test with Potassium Sulphate Solution the *USP* uses the salt itself and not the residue left on ignition, dissolves in Sulphuric Acid instead of Hydrochloric Acid, and uses a 1 in 10 Potassium Sulphate Solution in the place of a saturated solution. The *BP* makes no mention of the precipitates produced by Potassium or Sodium Hydroxide Solution.

It is officially required to lose 53 p c in weight when incinerated, but no direct indication is given of the composition of the ash. According to the *USP* it should leave a reddish-brown residue of Ceric and other rare earth oxides, amounting to not less than 47 p c of the salt. *BP* omits to mention that it should yield the reactions characteristic of Oxalates. Their presence may be detected on decomposing the salt by boiling with Potassium or Sodium Hydroxide.

Solution, adding a slight excess of Acetic Acid to the filtrate and then Calcium Chloride Solution, a white precipitate insoluble in Acetic Acid but soluble in Hydrochloric Acid should be produced

The more generally occurring impurities are Arsenic, Iron, Zinc, Calcium, Aluminium, Carbonates or Phosphates. These impurities are grouped collectively in the *B P*. The more important are Arsenic, which may be detected by the modified Gutzet's test, and Zinc, which may be detected by first boiling the salt with Potassium or Sodium Hydroxide Solution, cooling, filtering and adding Hydrogen Sulphide Solution to the filtrate. If this filtrate be previously divided into two portions, the second portion may be used in testing for Aluminium, which is precipitated on the addition of Ammonium Chloride Solution.

Iron may be detected by the addition of Potassium Ferrocyanide Solution to a solution of the salt in diluted Hydrochloric Acid. The *U S P* regards the absence of effervescence when the salt is dissolved in diluted Hydrochloric Acid as indicative of the absence of Carbonates, a test preferable to that of the *B P*, which requires that the reddish-brown powder remaining after decomposition at a dull red heat shall dissolve without effervescence in 'boiling Hydrochloric Acid'. Calcium and Phosphates are unlikely impurities and are not included in the *U S P*. Cerium Oxalate is not official in the *P G*.

Calcium Chloride—If the filtrate from the residue obtained by boiling the salt with Potassium Hydroxide T S be supersaturated with Acetic Acid, the addition of Calcium Chloride T S will produce a white precipitate insoluble in Acetic Acid, but soluble in Hydrochloric Acid, *U S P*.

Ammonium Carbonate—The solution in Hydrochloric or Sulphuric Acids yields with Ammonium Carbonate T S a white precipitate of Cerous and other rare earth carbonates, which are somewhat soluble in excess of the reagent, *U S P*.

Strychnine—If the residue left after heating Cerium Oxalate be dissolved in concentrated Sulphuric Acid, and a small crystal of Strychnine added, a deep blue colour will appear which will rapidly change to purple and then to red, *U S P*.

Time-limit Test—The Solution (1-20) in Diluted Hydrochloric Acid should not respond to the time limit test for heavy metals, omitting the addition of Ammonia Water, *U S P*.

Gutzet's Test—Five c.c. of a solution of the salt (1-10) in Diluted Hydrochloric Acid should not respond to the modified Gutzet's test for Arsenic, *U S P*.

Ammonium Chloride—On boiling the salt with T S of Potassium Hydroxide, and filtering, no precipitate should be produced in the filtrate by the addition of T S of Ammonium Chloride, *U S P*.

Ammonium Sulphide—The filtrate obtained on boiling the salt with T S of Potassium Hydroxide, and filtering, should not yield a precipitate with T S of Ammonium Sulphide, *U S P*.

CETACEUM

SPERMACEI

FR, BLANC DE BALEINE, GER, WALRAT, ITAL, CETINA,
SPAN, ESPERMA DE BALLENA

White, odourless, somewhat translucent, crystalline, pearly masses, unctuous to the touch, and having a bland mild taste. It

is a peculiar, concrete, fatty substance obtained from the Sperm Whale, *Physeter macrocephalus*, L., occurring chiefly in a cavity in the head, but also obtained from smaller cavities in the body

The Sperm Whale inhabits the Pacific and Indian Oceans

Cetin or Cetyl Palmitate when saponified yields Ethal (Cetyl Hydrate), and not Glycerin (Glyceryl Hydrate). Most Oils and Fats are Oleates, Palmitates, and Stearates of Glyceryl, which when saponified yield Glycerin and Oleates, Palmitates and Stearates of the metals

Solubility—Slightly in Alcohol (90 p c), 1 in 80 of boiling Alcohol (90 p c), 1 in 6 of Ether, 4 in 5 of Chloroform, and in the fixed and volatile Oils

Medicinal Properties.—Emollient It is much employed for ointments and cerates

Spermaceti can be powdered quite readily with the addition of Alcohol (90 p c), but when Alcohol is contra indicated it can also be powdered by the addition of 12 minims of fixed Oil of Almonds to each oz of Spermaceti

Official Preparations—Unguentum Cetacei Contained in Unguentum Aquae Rosae and Unguentum Capsici

Not Official—Mistura Cetacei, Cold Cream, Unguentum Cetacei sine Benzoino.

Foreign Pharmacopœias—Official in all except Belg. Mex. (Esperma), Port. (Espermaceti)

Descriptive Notes—Spermaceti is deposited from Sperm Oil, which is found in cavities of the head of the Sperm Whale, *Physeter macrocephalus*, L. The oil is cooled, the Spermaceti separated by settling, and pressed in bags to remove adhering oil, until hard and brittle. It is then melted in boiling Water to which a weak solution of Potash is added to remove impurities, washed and crystallised. The crystalline masses in which it occurs in commerce are quite characteristic in appearance. Chinese insect wax, which is but rarely imported, closely resembles it, but is much harder. It is a wax secreted by *Coccus Pera*, Westw., in China, on the twigs of Chinese species of Ash and Privet, and is used by the Chinese to form an outer layer on candles to prevent guttering. It has a much higher melting point than Spermaceti.

Tests.—The distinguishing tests for Cetaceum are the melting point, which should be about 48° C (118 4° F) and which is officially required to be from 46° to 50° C (114 8° to 122° F), but which is occasionally as low as 43° C (109 4° F), and the specific gravity, which should be from 0 942 to 0 946, and to which no reference is made in the B.P. The U.S.P. gives the melting point as 42° to 50° C (107 6° to 122° F), the P.G. 45° to 50° C (113° to 122° F). The B.P. adopts the same method for taking the melting point of Cetaceum as for Cera Flava. The U.S.P. gives the specific gravity as 0 935 to 0 944 at 25° C (77° F), 0 842 at 100° C (212° F), the P.G. gives an average specific gravity of 0 943. It should possess an Acid value of from 0 1 to 0 5, and a Saponification value of from 125 to 136. The B.P. places the limit of acidity at not more than 'one drop' of Normal Volumetric Sodium Hydroxide Solution

for the neutralisation of the alcoholic solution of 0.2 gramme of Spermaceti, using Phenolphthalein Solution as indicator, but gives no figure for the Ester value. The quantity (0.2 gramme) used in the official method of determination is too small, and it is preferable to employ from 1 to 5 grammes. The limit of acidity indicates an Acid value of 13.9, and is altogether too high even for old samples. The Acid value increases greatly with age. A sample examined in the author's laboratory in November, 1896, showed the merest trace of free acid, but when re-examined in July, 1901, exhibited an Acid value of 4.5. The Iodine absorption of pure Spermaceti is practically nil.

The more generally occurring sophistications are Stearic and Palmitic Acids, Stearin, Tallow, and Paraffin Wax. Stearin and Palmitic Acid may be detected by the reaction of the alcoholic solution towards Litmus and by the precipitate produced on dilution of the solution with Water. The *USP* and *PG* boil with anhydrous Sodium Carbonate and Alcohol, subsequently acidifying the solution with Acetic Acid, and state that the solution may become turbid, but should not give a precipitate. The solubility in boiling Alcohol (90 p.c.) detects the presence of Stearin, Tallow, and Paraffin Wax. Tallow and Stearin increase and Paraffin Wax diminishes the saponification value. The presence of Tallow may also be recognised by the increased Iodine absorption. Paraffin Wax notably diminishes the specific gravity.

Alcohol.—If Spermaceti be boiled with Alcohol (90 p.c.), and the mixture cooled and filtered, the filtrate should not afford a flocculent precipitate on the addition of Water, *BP* (an equal quantity of Water), *PG*. At ordinary temperatures Cetaceum gradually crystallises out from a solution in boiling Alcohol (90 p.c.), which is approximately 1 in 50, *PG*.

Acetic Acid.—If 1 gramme of Cetaceum be boiled with 1 gramme of anhydrous Sodium Carbonate and 50 c.c. of Alcohol, and the mixture cooled and filtered, the filtrate on being supersaturated with Acetic Acid may become turbid, but it should not afford a precipitate, *PG* and *USP*.

Preparations

UNGUENTUM CETACEI SPERMACEI OINTMENT

Spermaceti, 5, White Beeswax, 2, Almond Oil (by weight), 18, Benzoin, in coarse powder, 1. Melt together the Spermaceti, Beeswax, and Almond Oil, add the Benzoin, and frequently stirring the mixture, continue the application of heat for two hours, remove from the source of heat, strain, and stir the Ointment constantly until cold.

It would be better to omit the Benzoin, which was first added in 1865, as pointed out in previous editions, the Benzoin converts this emollient preparation into one which is irritating: see below.

The following are called **Unguentum Cetacei**—(all by weight).

Dan. Spermaceti 2, White Wax 1, Oil of Almonds 12, Rose Water 5.
Norw. Spermaceti 6, White Wax 6, Oil of Almonds 58, Rose Water 30.
Russ. Spermaceti 3, White Wax 3, Olive Oil 14, Rose Water 2.
Swed. Spermaceti 2, White Wax 1, Oil of Almonds 12, Rose Water 5.
Swiss. Spermaceti 2, White Wax 1, Almonds Oil 7, Ethereal Tincture of Benzoin, 1.

The following are called **Ceratum Cetacei**—(all by weight) —

Austr Spermaceti, White Wax, Sesame Oil, equal parts

Hung Spermaceti 8, White Wax 8, Lard 9

Port Spermaceti 1, White Wax 1, Oil of Almonds 3

Span Spermaceti 4, White Wax 4, Oil of Almonds 47, Rose Water 45

U S See **Unguentum Aquæ Rosæ**

Cold Cream is a synonym for Unguentum Cetacei, *Dan* and *Swed*, and Unguentum Leniens, *Belg* and *Ger*

The following are called **Unguentum Leniens**—(all by weight) —

Austr Spermaceti 15, White Wax 8, Sesame Oil 62, Water 15, Otto of Rose 2 drops

Belg White Wax 14, Almond Oil 56, Rose Water 30

Dutch Spermaceti 10, Yellow Wax 5, Adeps Lanae 10, Sesame Oil 50, Rose Water 25

Ger Spermaceti 8, White Wax 7, Almond Oil 57, Water 28, Otto of Rose 0 1

Unguentum Refrigerans—(all by weight) —

Swiss Spermaceti 10, White Wax 8, Arachis Oil 57, Castor Oil 5, Otto of Rose 1 drop, Rose Water 20

Pomata con Olio di Mandorle —*Ital* Spermaceti 1, White Wax 1, Oil of Almonds (by weight) 8

Cerato de Galeno —*Span* Almond Oil 55, White Wax 15, Agua de Rosas 30

Not Official

UNGUENTUM CETACEI SINE BENZOINO —Spermaceti 5, White Beeswax 2, Almond Oil 18

The *B P* ointment made with Benzoin is unsuited for many purposes for which this ointment is useful, such as eye ointments, &c.

Used as a cooling dressing Applied on lint to broken blisters &c. After walking, it affords great relief, and frequently enables one to continue the exercise without serious discomfort It is also recommended for smearing on the feet before starting for a long walk on rough ground

MISTURA CETACEI —Spermaceti 60 grains, Proof Spirit 15 minims, finely pulverise the Spermaceti by aid of the Spirit, and add by degrees half the yolk of an egg, at first only sufficient to make a stiff paste, which should be made very smooth by diligent trituration, then add the rest, and make up with Water to 4 oz

[This formula was given in Squire's *Companion* 1864]

COLD CREAM —White Beeswax, 1, Spermaceti, 1, Oil of Almonds, 8, Rose Water, 11, Otto of Rose to perfume it Melt together, by means of water-bath, the Oil, Spermaceti, and Beeswax, add the Otto, strain through muslin into the Rose Water, stir together whilst gently warming until Water globules are no longer visible, and the mixture is of proper consistence to pour into pots without separating This form has been used by the author for several years, but Cold Cream is more easily and more generally made with less Rose Water See also Unguentum Aquæ Rosæ under Rosæ Oleum

Cold Cream —Spermaceti, 60, White Wax, 30, Almond Oil, 215; Rose Water, 60, Tincture of Benzoin, 15, Otto of Rose, 10 drops —*Fr*

Not Official.

CETRARIA

ICELAND MOSS

The dried Lichen, *Cetraria Islandica*, L. A native of the north of Europe

It contains a bitter principle, **Cetrarin** (Cetraric Acid), which has been used as a tonic.

Medicinal Properties —Demulcent, nutritious, and slightly tonic

Iceland Moss Jujubes are useful for coughs

Foreign Pharmacopœias —Official in Austr, Belg, Dutch, Fl (Lichen d'Islande), Ger, Hung, Ital, Jap, Port, Span (Liquen Islandico), and Swiss, *Lichen Islandicus*, Mex (Liquen de Islandia) Not in Norw

Descriptive Notes —The frond of the lichen, *Cetraria islandica*, L., is more or less branched in a forked manner, of a brownish colour above, and greyish white below, marked on the under surface with numerous minute scattered chalky white pits. It is flat, channelled above, and has a crisp mode of growth. The wavy edges are furnished with a fringe of numerous minute short linear papillæ. The fructification, which consists of a flat, disc like, dark brown expansion near the margin of the frond, is rarely met with, and then mostly on the broader varieties of the plant. The taste is mucilaginous and slightly bitter.

DECOCTUM CETRARIÆ —Iceland Moss, 1, first wash with cold Water, then add Distilled Water, 20, boil ten minutes, strain with gentle pressure whilst hot and wash the marc to make 20 (1 in 20)

Dose —1 to 4 fl oz = 28.4 to 113.6 grammes

Official in Ital, 1 in 20, Fr (Tisane), 1 in 100, Ital has also **Infusion**, 1 in 20

SACCHARUM CETRARIÆ —Iceland Moss 1, Sugar 1, Water 100. Wash the Iceland Moss with Water to remove the bitterness, then boil with 100 of Water, strain and express lightly, and in the strained liquid dissolve the Sugar and evaporate on a water-bath. When sufficiently firm remove from the bath and dry in a cupboard to a powder or scale.

GELATINA CETRARIÆ (Iceland Moss Jelly) —Saccharated Cetraria 2 Sugar 1, Water 5. Mix, boil gently till scum collects on the surface, then withdraw the heat, remove the scum, and pour into pots to cool.

A similar preparation is given in Ital, Port and Span.

Paté de Lichen officinal containing 1 of Extract of Opium in 5000 —Fr

Not Official

CARRAGEEN

IRISH MOSS *Syn* —CHONDRUS

Fr, CARRAGAHEEN, GER, IRLANDISCHES MOOS, ITAL, FUCO CARAGEO, SPAN, CARRAGAEN

The dried seaweed *Chondrus crispus*, Lyngb. It is used as an article of food on the west coast of Ireland, where it abounds. Has been proposed as a substitute for Acacia as an emulsifying agent and for the suspension of some powders.

One part of *Chondrus* boiled for ten minutes with 30 parts of Water yields a solution which gelatinises on cooling, and is not coloured blue by T.S. of Iodine —U.S.P.

Foreign Pharmacopœias —Official in Belg, Dan, Dutch, Fl, Ger, Hung, Ital (Fuco Carageo), Jap, Mex (Liquen Carragaen), Port (Alga Perlada), Span and Swiss. Fl has a Tisane 1 in 200.

SACCHARUM CARRAGEEN —Made like Saccharum Cetrariæ

GELATINA CARRAGEEN (Irish Moss Jelly) —Made like Gelatina Cetrariæ

Official in Port

MUCILAGE OF MOSS —Irish Moss, $\frac{1}{2}$ oz, is boiled with 40 oz of Water for 15 minutes and made up to 34 oz —*Arnold's Formulary*

Decoction Chondri —Irish Moss, 2 50, Distilled Water, q s to produce 100 —B.P.C.

The B.P.C. Supplement gives the *syn*. Mucilage of Irish Moss

Not Official

CHALMOOGRA OIL *See* GYNOCARDIÆ OLEUM

Not Official
CHELIDONIUM.

GREATER CELANDINE

The entire Plant *Chelidonium majus*, L

The juice has been used in opacities of the cornea, and is a popular application for the cure of warts — *B M J* '97, i 25 and 354

Has been recommended chiefly by Demisonko in the treatment of cancer: *B M J* '97, i 25, 354 and 637, ii 123, *B M J E* '96, ii 88, '97 ii 47, *L* '96, ii, 649 and 1778, *L* '97, ii 737, *P J* '97, i 86 Unfavourably commented on *L J* '98, i 61. In the treatment of inoperable cancer, Celandine is worthy of trial — *L* '01, ii 967, *C D* '01, ii 1048

Chelidonium — This alkaloid forms colourless crystals, melting at 135° C (275° F), soluble in Alcohol (90 p c), insoluble in Water, and but slightly soluble in Ether

Dose — 1 to 3 grains = 0.06 to 0.2 gramme

The **Sulphate** is readily soluble in Water, the **Hydrochloride** less so, and the **Tannate** is insoluble in Water

CHIRATA.

CHIRETTA

FR, CHIRETTE, GER, OSTINDISCHER ENZIAN

The entire dried Plant, *Swertia Chirata*, Ham, collected when in flower

It is a native of, and is obtained from, Northern India

Under the title of **Andrographis**, the dried *Andrographis paniculata*, is official in the *Ind* and *Col Add* for India and Ceylon

Medicinal Properties — Bitter tonic and stomachic, without astringency, given in atonic dyspepsia. Containing no Tannin, it may be prescribed with Iron

Official Preparations — Infusum Chiratae, Liquor Chiratae Concentratus, and Tinctura Chiratae

Not Official — Infusum Chiratae Concentratum

Foreign Pharmacopœias — Official in Port and U S. Not in the others.

Descriptive Notes — The Chirata official in the *B P* is distinguished by its very bitter taste, by the opposite, entire, glabrous, ovate leaves, and by the pith being continuous, solid, and easily separable. It is met with in commerce in compact flattened bundles about 3 ft long and about 5 or 6 in thick, and weighing 1½ to 2 lb, and bound round with a slip of bamboo. Other allied species are used in different provinces of India, and some of them are occasionally imported and sold as the genuine drug. Of these *Swertia* *parviflora*, B & H, has a more woody, tougher and nearly hollow stem. *S. pinnatifida*, R & S, has pale angular stems with a large pith. These give a much less bitter infusion than genuine Chirata. Sometimes Chirata is falsely packed, Madder roots or other plants, or even stones, being concealed in the centre of the bundle. *Andrographis parviflora*, Nees, has distinctly quadrangular stems and irregular flowers, and only

3 or 4 seeds in each capsule, and is not packed in loose bundles. It is rarely imported and not likely to be confounded with *Chnata*. It is official in the *Ind* and *Col Add*. Under the microscope the leaves may be distinguished from those of *Chnata* by the presence of cystoliths below the upper epidermis, which are shorter than in most other medicinal Acanthaceous plants, by the stomata being placed between a large and small cell, and by the quadricellular glandular hairs (*Planchon* and *Collin*)

Preparations

INFUSUM CHIRATÆ

Chiretta, cut small, 1, boiling Distilled Water, 20 infuse for fifteen minutes, strain (1 in 20)

Now 1 in 20 instead of 1 in 40

Dose — $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

A corresponding preparation, *Infusum Andrographidis*, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

LIQUOR CHIRATÆ CONCENTRATUS CONCENTRATED SOLUTION OF CHIRETTA

1 of Chiretta, in No 40 powder, percolated with Alcohol (20 p c) to yield 2 (1 in 2)

Dose — $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

Tests —Concentrated solution of Chiretta has a specific gravity of 0 990 to 1 010, it contains about 5 p c w/v of total solids and 19 p c w/v of Absolute Alcohol

A corresponding preparation, *Liquor Andrographidis Concentratus*, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

U S has a **Fluid extract**, 1 in 1, using Alcohol 49 p c

TINCTURA CHIRATÆ

1 of Chiretta, in No 40 powder, percolated with Alcohol (60 p c) to yield 10 (1 in 10)

Dose — $\frac{1}{2}$ to 1 fl drm = 1 8 to 3 6 c c

Prescribed in 5 minim doses, with Acids and Tincture of Orange to form an acid tonic mixture

A corresponding preparation, *Tinctura Andrographidis*, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Tests —Tincture of Chiretta has a specific gravity of 0 920 to 0 925, it contains about 1 5 p c w/v of total solids and about 60 p c w/v of Absolute Alcohol

Not Official

INFUSUM CHIRATÆ CONCENTRATUM —Chiretta, in No 40 powder, 40, Alcohol (90 p c), 25, Dilute Chloroform Water (1 in 1000), sufficient to make 100. Prepare by repercolation —*Fun* and *Weight*, P J '06, 1 165 and '07, 1 621, C D '06, 1 252, Y B P 1907, 249

This formula appears in the B P C

CHLORAL HYDRAS.

CHLORAL HYDRATE

CCl, CH(OH)₂, eq 164 15

FR, HYDRATE DE CHLORAL, GER, CHLORALUM HYDRATUM, ITAL, CLORALIO IDRATO, SPAN, HIDRATO DE CLORAL

Transparent, colourless, rhomboidal crystals, having an aromatic penetrating odour, and an unpleasant, slightly bitter, acrid taste

It should be kept in well-stoppered amber-tinted glass bottles in a cool and dark place

Chemically it is Trichlorethylidene Glycol Anhydrous Chloral is produced by the action of dry Chlorine gas on Ethyl Alcohol, the pure Chloral being subsequently converted into Hydrate by the addition of the necessary amount of Water and purified by recrystallisation from suitable solvents

Solubility.—4 in 1 of Water, and measures $3\frac{1}{4}$, 5 in 1 of Alcohol (90 p c), 2 in 1 of Ether, 2 in 1 of Glycerin, 1 in 1 of Olive Oil, 1 in 3 of Chloroform, 1 in 10 of Oil of Turpentine (cold), 1 in 5 boiling, 1 in 68 of Carbon Bisulphide

Medicinal Properties—An excellent hypnotic, producing natural and placid sleep soon after its administration, in acute mania and delirium tremens it is given as a cerebral depressant Given in asthma and $\text{v' } \text{u' } \text{c' }$ and extreme cases of chorea, efficacious in large doses in sea-sickness Has been found useful as a spinal depressant and antispasmodic in tetanus, uræmic and puerperal convulsions, and by intravenous injection in Strychnine poisoning Of great value in labour, as it relieves pain, assists to dilate the os and relax the rigid perinæum, especially in primiparæ, without lessening the expulsive power of the uterus Given in nocturnal incontinence of urine It should not be given in advanced cardiac disease, nor in fatty heart Children stand it well

It is not suitable for insomnia due to pain, as an analgesic it is inferior to opium

In concentrated solution, applied locally, it acts as a vesicant

As a pungent irritant with Camphor and sometimes with Cocaine, it is useful for the relief of neuralgia, rheumatism, toothache and chilblains

In tetanus in 20 grain doses every five hours (*B M J* '04, ii 1460), and along with injections of antitetanic serum (*B M J* '04, ii 1429), in sea-sickness teaspoonful doses every five minutes of a mixture of 2 drms of the syrup with $\frac{1}{2}$ drm Ammonium Bromide made up to 1½ oz with Water, where this fails relief often follows 1 minim doses of Tincture of Iodine — *B M J* '04, ii 1405

The only satisfactory way of giving this drug in obstinate cases of sea sickness is by the rectum in doses of from 20 to 30 grains — *B M J* '05, i 1090

The tendency to eclampsia may often be controlled (*B M J* '05, ii 718) by the administration of $\frac{1}{2}$ drm doses, combined with a drm of Potassium Bromide and repeated if necessary every hour

It is shown (*B M J* '05, ii 250) to be of special use in sleeplessness and pain of gouty people with high blood pressure It seems a general law (*B M J* '05, ii 1005) that when a hypnotic contains Chlorine in its molecule its effects are not limited to the brain and central nervous system, but extend to the heart and the tissues in general

In the treatment of convulsions in early infancy (*Pr* lxxv 514), 2 to 3 grains may be given hypodermically or introduced into the bowel through a rubber catheter. For rectal injection 5 grains may be given to a baby of six months and 10 grains to one a year old. One grain every two hours in the youngest babies (*Pr* lxxv 517), and in children of one or two months 1 to 2 grains, continued in these doses until the fits have ceased, for at least 24 or 36 hours, in the treatment of idiopathic convulsions.

Effects from an overdose or repeated overdoses are excitement, convulsions, and delirium, followed by deep coma and quiet sleep from which the patient may never stir, he may, however, pass to death without any previous convulsions. It lowers temperature, and causes contraction of the pupil.

A case of puerperal eclampsia treated by Chloral Hydrate, Potassium Bromide, and Chloroform inhalation — *L* '97, ii 915

As a pigment to the interior of the nostrils in acute coryza, 10 grains in 4 drms Castor Oil — *Pr* lv 517

Applied spread over the surface of diachylon plaster, the skin having been previously rubbed over with Almond Oil or Vaseline, it acts as a vesicant, superior to Cantharides — *P J* '02, i 115

Dose — 5 to 20 grains = 0.32 to 1.3 gramme

Ph Ger maximum single dose, 3 grammes, maximum daily dose, 6 grammes

Prescribing Notes — 3 oz will dissolve in 1 fl oz of Water, and measure 2 fl oz and 5½ fl drms, if to this be added 23 minims of Water, every minim will contain a grain of Chloral. This solution is handy for dispensing.

It is usually given in solution, but the objectionable taste is difficult to mask, Chloral Hydrate, 20 grains, Syrup of Orange, 1 fl drms, Peppermint Water, to 1 fl oz, make a good draught for those who do not object to Peppermint.

Chloral Hydrate, 4 grains, Liquorice Root, in powder, 1 grain, Gum Acacia, in powder, ½ grain, make a good pill, with a trace of 'Diluted Glucose'.

The addition of 1 grain to the fl oz will keep hypodermic solutions otherwise liable to develop fungoid growths.

Incompatibles — When prescribed with Alkalis, Chloroform will be liberated.

Official Preparation — Syrupus Chloral

Not Official — Liquor Bromo Chloral Compositus, Chloral Camphoratum, Chloral cum Camphora et Cocaina, Chloral et Phenol, Suppositoria Chloral, Dormiol, and Chloral Tannum.

Antidotes — Stomach tube or emetics, keep up the temperature by hot blankets, hot water bottles, etc., injection of a pint of hot strong coffee into rectum, electric magnetism, inhalations of Amyl Nitrite, in bad cases hypodermic injection ½ grain of Strychnine Nitrate, artificial respiration — *Murrell* ½ of a grain of Picrotoxin has been found enough for 30 grains of Chloral — *B M J* '75, i 506

Foreign Pharmacopœias — Official in Austr, Belg, Ger, Hung, Jap, Russ, Swiss and US (Chloralium Hydratum), Dutch (Hydras Chlorali), Dan, Norw and Swed (Hydras Chloralicus), Fr, Ital (Clorale Idrato), Mex (Cloral Hidratado), Port (Hydrato de Chloral), Span (Hidratado de Cloral).

Tests — The distinguishing tests for Chloral Hydrate are the melting point of the crystals, which, when dried, should be about 58° C (136 4° F), the temperature at which the melted liquid again becomes solid (the solidifying point), which is officially required to be about 48 9° C (120° F), and the boiling point (when heated in a test-tube with pieces of glass contained in it), which is officially required to be from 94 4° to 96 7° C. (202° to 206° F). The *B P*

gives no figure for the melting point, but *PG* and *USP* state it to be about 58°C (136.4°F) when dry. The *USP* gives a range of from 35° to 50°C (95° to 122°F) for the solidifying point. The best solidifying point is considered by *USP* to be 122°C (252°F). If the sample is too much under-hydrated the solidifying point is lower and the boiling point is under 95°C (203°F), and the sample is prone to decompose and become acid on keeping. If over-hydrated the solidifying point is lower and the boiling point is above 98°C (208.4°F) and the sample is deliquescent, a slightly under-hydrated sample is the best for good-keeping qualities.

The aqueous solution should be neutral or but slightly acid to Litmus. The *USP* states that the aqueous solution gradually becomes an acid reaction, but that a neutral alcoholic solution remains permanently neutral.

An aqueous solution warmed with Potassium or Sodium Hydroxide Solution evolves immediately a powerful odour of Chloroform, the Chloral Hydrate being decomposed with formation of Chloroform and the Formate of the alkali Hydroxide. If to the cooled solution a few drops of Aniline Oil be added, and the liquid again warmed, the powerful, penetrating, and highly poisonous odour of Phenyl-isocyanide is evolved. When heated with an excess of Potassium or Sodium Hydroxide Solution the Chloroform at first formed is itself decomposed with the production of the corresponding Chloride of the alkali and a further quantity of Formate.

The *BP* process of quantitative determination is founded upon decomposition by an alkali Hydroxide Solution. It is officially required to indicate 98.5 p.c. of pure Chloral Hydrate as volumetrically determined from the quantity of Normal Volumetric Sodium Hydroxide Solution required to decompose 4 grammes. No requisite percentage or method of determination is given in the *PG*. The *BP* test has been severely criticised. If as in the official method of procedure the Chloral Hydrate be 'heated' with the Normal Volumetric Sodium Hydroxide Solution, results greatly in excess of the truth are yielded, attributable to secondary decomposition of the Chloroform by the excess of alkali. The term 'heated' may be variously understood by different operators. It has been pointed out (*PJ* '99, i 236, '01, i 387, '03, i 531) that if the reaction be allowed to proceed at the ordinary temperature fairly concordant results may be obtained, but the range of temperature must be restricted within narrow limits. A correction may also be made (*PJ* '07, ii 4) by allowing the secondary reaction to commence and ascertaining the extent to which it has proceeded by titration of the resultant Chloride with Deci-normal Volumetric Silver Nitrate Solution.

The *Companion* (17th Edition) suggested determination, when in very dilute solution, by reduction by the Copper-Zinc couple and titration with Deci-normal Volumetric Silver Nitrate Solution. A process described (*PJ* '07, ii 6) depends upon the reduction of the Chloral Hydrate by means of Aluminium Powder or Zinc filings and Acetic Acid. A weighed quantity of 0.25 gramme is boiled with 1 gramme of Aluminium Powder, 15 c.c. of Acetic Acid *BP*, and

40 c.c. of Water under a reflux condenser for half an hour. The mixture is then filtered, the filter and flask washed with Water, 50 c.c. of Deci-normal Volumetric Silver Nitrate Solution is then added, the Silver Chloride filtered out, and the excess of Volumetric Silver Nitrate Solution determined with Deci-normal Volumetric Ammonium Sulphocyanide Solution, adding 10 c.c. of strong Nitric Acid and 5 c.c. of a saturated Iron Alum solution as indicator. Each c.c. of Deci-normal Volumetric Silver Nitrate Solution used is equivalent to 0.005472 gramme of pure Chloral Hydrate.

Chloral Hydrate may be extracted from its aqueous solution by shaking out with Ether or Acetic Ether.

A simple process for the approximate determination of the amount of Chloroform yielded on treatment with Potassium Hydroxide may be conveniently conducted in a graduated tube, thus: Place in a tube 250 grain-measures of a 20 p.c. Potassium Hydroxide Solution, and add to it gradually (keeping it cold) 50 grains of the Chloral Hydrate, cork securely, and shake, allow the liquid to separate, and the number of grain-measures of Chloroform (at the bottom), to which must be added 1 for every 200 grain-measures of supernatant liquid, multiplied by 1.5 gives the grains of Chloroform, which should be not less than 35.

The more generally occurring impurities are mineral matter, Chloral Alcoholate, certain organic impurities and Chlorides. Mineral residue is readily detected when the drug is volatilised on Platinum foil. The *B.P.* employs the Iodoform test as a means of detecting Chloral Alcoholate. No yellow crystalline precipitate of Iodoform should be produced within an hour, when Iodine Solution, in sufficient quantity to yield a brown coloration, is added to a filtered mixture of 1 gramme of the hydrate warmed with 6 c.c. of Water and 0.5 c.c. of Potassium Hydroxide Solution, *B.P.* Organic impurities may be detected by shaking the chloroformic solution with concentrated Sulphuric Acid, whilst Hydrochloric Acid and Chlorides may be detected by Silver Nitrate Solution, the *B.P.* requires that the aqueous solution should not afford any precipitate with Silver Nitrate Solution, the *U.S.P.* requires that the 1 in 20 aqueous solution, slightly acidulated with Nitric Acid, should remain unaffected by Silver Nitrate T.S.

Sulphuric Acid—No colour should be imparted to Sulphuric Acid when it is shaken with a solution of Chloral Hydrate in Chloroform, *B.P.* The *P.G.* test directs 0.5 gramme of Chloral Hydrate to be vigorously shaken with 5 c.c. of Sulphuric Acid in a glass-stoppered test-glass of 3 cm. diameter, and which has been previously rinsed out with Sulphuric Acid. The Sulphuric Acid should not become coloured within one hour.

Silver Nitrate—An alcoholic solution (1-10) of Chloral Hydrate should not be affected at once by T.S. of Silver Nitrate, *P.G.*, that an aqueous solution (1-20) slightly acidulated with Nitric Acid should remain unaffected, *U.S.P.*, an aqueous solution should not afford any precipitate with T.S. of Silver Nitrate, *B.P.*

Preparation

SYRUPUS CHLORAL SYRUP OF CHLORAL

Dissolve 800 grains of Chloral Hydrate in 15 fl. drms. of Distilled Water, and add Syrup, *q.s.* to yield 10 fl. oz. (10 grains in 60 minims)

Dose.— $\frac{1}{2}$ to 2 fl drms = 18 to 71 cc

Foreign Pharmacopœias—Official in Belg and Fr, 1 in 20, with Peppermint, Mex, 1 in 20, Port, 1 in 50, Span, 1 in 32.5 All by weight Not in the others

Not Official

LIQUOR BROMO-CHLORAL COMPOSITUS—Chloral Hydrate, 1600 grains, Tincture of Indian Hemp, 400 minims, Tincture of Orange, 400 minims, Henbane Juice, 1600 minims, Syrup, $3\frac{1}{2}$ fl oz, Fluid Extract of Liquorice, $\frac{1}{2}$ fl oz, dissolve. Add 1600 grains of Potassium Bromide dissolved in 7 fl oz of Distilled Water, filter, wash with Distilled Water to produce 20 fl oz. Each fl drim contains 10 grains of Chloral Hydrate and 10 grains of Potassium Bromide

Dose.— $\frac{1}{2}$ to 2 fl drms = 18 to 71 cc—*BPC Formulary* 1901

BPC is practically the same strength, but calculated into parts per 100 fluid parts

Bromidia is somewhat similar in composition

It has been suggested that each fl drim should be made to contain 15 grains each of Chloral and Potassium Bromide, and that the filtration should be omitted, since it takes out the resins of the Indian Hemp. Suspension by mucilage recommended. Hyoscine Hydrobromate recommended to be substituted for Henbane—*CD '02*, n 314

SUPPOSITORIA CHLORAL—Chloral Hydrate, 5 grains, White Wax, 5 grains, Oil of Theobroma, 7 grains. Melt together the Wax and Theobroma Oil, and when partially cooled, mix in the Chloral Hydrate and pour into a mould

CHLORAL CAMPHORATUM—Chloral Hydrate, 1, Camphor, 1, rub together in a warm mortar until completely liquefied, and filter if necessary—*BPC Formulary* 1901, now incorporated in the *BPC*

As a **Pigmentum** this formula has appeared for many years in the Pharmacopœias of the London, Throat and Westminster Hospitals

Useful application for the relief of neuralgia

CHLORAL CAMPHORATUM CUM COCAINA—Chloral Hydrate, 9, Camphor, 9, Cocaine, 2

This has been incorporated in the *BPC*

CHLORAL ET PHENOL—Chloral Hydrate, 1, Carbolic Acid, 1

Is soluble in Water, Alcohol (90 p c), and in Glycerin

The Chloral Carbolatum of *BPC*

So long as the proportion of Carbolic Acid to Chloral does not exceed 1.7 to 1, the product will mix with Water in all proportions, beyond this limit the excess of Carbolic Acid separates on the addition of Water. As it corresponds to 8 molecular weights to 1, there is probably a chemical combination in these proportions—*PJ (3) xvi* 188

DORMIOL (Amylene Chloral)—A colourless liquid possessing a camphoraceous odour, usually supplied commercially as a 50 p c solution. mixes readily with Water. It is also supplied in capsules.

A good narcotic in mental diseases, and stated to produce no untoward effects—*L '99*, n 73, '02, n 1712, *BMJ '02*, n 1278, *PJ '03*, n 62

Dose.—5 to 20 minims = 0.3 to 1.2 cc

Prescribing Notes—It can be given in capsules, or in mixtures, covering the unpleasant taste with Syrup of Tolu, using equal parts of the Syrup and Water

CHLORAL TANNIN (Captol).—A brown, resinous substance, soluble in Water and in Alcohol

A solution has been introduced as a Hair Wash.

Not Official

CHLORALAMIDUM

CHLORALAMIDE CHLORALFORMAMIDUM, U S , CHLORAL FORMAMIDE

$\text{C}_3\text{H}_4\text{Cl}_2\text{NO}_2$, eq 191 00

Chloralamide is a compound of Chloral Anhydride and Formamide Colourless lustrous, odourless crystals, possessing a somewhat bitter taste Its aqueous solution should not be heated above $48\text{ }^\circ\text{C}$ ($120\text{ }^\circ\text{F}$), as above that temperature it undergoes hydrolysis, being reconverted into Anhydrous Chloral and Formamide It is permanent in weakly acidulated solutions, but decomposed by alkalis

It should be preserved in well stoppered amber tinted glass bottles

Care should be taken not to confound Chloralamide with Chloralimide

Solubility—1 in 21 of Water, 1 in 2 of Alcohol (90 p c), it dissolves very slowly in Glycerin but if sufficient time is allowed, 1 in 12 solution can be obtained, in about 3 days at $60\text{ }^\circ\text{F}$

Published solubilities of it in Water have varied considerably The *Companion* figure (1890) as above has been confirmed (*P J* (3) xxii 805), with the additional note that below $60\text{ }^\circ\text{F}$ the solubility decreases very rapidly —*C D* '92, 1 445

Medicinal Properties—Hypnotic It is stated to have much less influence on the heart than Chloral, and therefore may be used in cardiac disease, and that the dose need not be increased after continued use

Given in all kinds of insomnia —*L* '89, ii 849, 1192, '90, 1 339, *B M J* '89, ii 1326 '91, 1 1060, *M P* '89, ii 571, *P J* (3) xxi 104, *T G* 91, 634, 757, *Pr* xlvii 274 *B M J* '05, ii, 1007 In insomnia with 'irregular' heart after influenza —*B M J* '94, ii 1045

The safest of the hypnotics for the insomnia of cardiac disease —*B M J* '97, ii 857 '05, ii 250 Chloralamide is safer but slower in action than Chloral Hydrate —*L* '99, ii 143 20 to 30 grains in a little spirit, useful as a sleeping draught for patients suffering from acute Bright's disease —*Pr* lxxvii 645 658

In the insomnia of enteric fever—30 to 40 grains, repeated in lesser amount in two hours —*B M J* '04, ii 1452

Prescribed with Potassium Bromide as a remedy for sea sickness —*Pr* lvi, 145

Dose—20 to 50 grains = 1 3 to 3 24 grammes

Ph Ger maximum single dose, 4 grammes, maximum daily dose, 8 grammes

Prescribing Notes—It is prescribed in aqueous mixtures suspended with *Tragacanth*, or dissolved in weak Alcohol or Glycerin, see *Haustus* and *Mistura* given below

Foreign Pharmacopœias—Official in Ger (Chloralium formamidum), Mex (Chloralamido), U S (Chloralformamidum) Not in the others

Tests—Chloralamide has a melting point of 114 ° to $115\text{ }^\circ\text{C}$ (237 ° to $239\text{ }^\circ\text{F}$) The aqueous solution of the salt should be neutral in reaction towards Litmus paper It yields a turbid solution when warmed with Potassium or Sodium Hydroxide Solution, the solution clearing on the separation of the Chloroform produced by the decomposition, the evolved vapour possessing an alkaline reaction towards red Litmus paper

The more generally occurring impurities are inorganic matter, Chloral Alcoholate, Ethyl Carbamate, free acids, *e g*, Formic and Hydrochloric Acids, and products of decomposition Inorganic matter, Chloral Alcoholate, and Ethyl Carbamate are detected by the behaviour of the sample when carefully heated in an open dish no weighable residue should remain after ignition, and no inflammable vapours should be given off Free acids are detected by the reaction of the alcoholic solution towards moistened blue Litmus paper, and products of decomposition by Silver Nitrate Solution, no immediate turbidity should be produced on the addition of a few drops of Silver Nitrate Solution to a 10 p c alcoholic solution

Not Official

CHLORETONETRI CHLOR TERTIARY BUTYL ALCOHOL CHLOR-BUTYL ALCOHOL
ACETONE CHLOROFORM $C_7H_9Cl_3$ (OH) eq 176.09

Light, white, glistening crystals, having a strong camphoraceous odour and taste. It is volatile at the ordinary temperature of the air.

It may be obtained by the interaction of Chloroform, Acetone, and an alkali Hydroxide.

This must not be confused with Chloroform prepared from Acetone, which is also known as 'Acetone Chloroform'.

Solubility—1 in 125 of Water, 6 in 4 of Alcohol (90 p.c.), also soluble in Chloroform and Ether, readily soluble in Glycerin and in Clove Oil.

Introduced as a hypnotic and local anæsthetic. It is also stated to possess slight analgesic and antiseptic properties.—*T'G* xxiv 18, 98, *L* '00, 1 106.

In epilepsy.—*T'G* '01, 757.

As a hypnotic in 25 cases of mental disease, 1 to 1½ gramme doses. In restless subjects, 2 grammes may be given.—*B M J E* '02, 1 31.

The therapeutic and the toxic doses (*B M J E* '05, 11 4) are so near one another that this drug has been discarded as a hypnotic altogether.

To prevent post operative vomiting.—*P J* '03, 1 340.

5 grain doses every three hours to prevent sea sickness.—*L* '03, 1 615, 687, *C D* '03, 1 424.

5 grains Chloretone 15 minutes before embarking will generally ensure complete immunity during the Channel passage even in rough weather, and for longer voyages in rough weather it should be taken 20 minutes before meals two or three times a day, an efficient remedy in cholera, has a remarkably soothing effect in painful and irritable stomach conditions, a hypnotic in 10 grain doses (*L* '07, 1 880), skin eruption due to chloretone treatment of cholera.—*L* '07, 1 883.

Dose—5 to 20 grains=0.32 to 1.3 grammes.

Prescribing Notes—Conveniently given when dissolved in a mixture of Alcohol and Glycerin. It is suspended with difficulty by Mucilage of Gum Acacia or Tragacanth. Cachets and Powders should be enclosed in a bottle, but even under these circumstances there is considerable loss in a month or two.

Tests—Chloretone melts at 80° to 81° C (176° to 177.8° F), but when anhydrous the melting point is raised to 95° to 96° C (203° to 204.8° F). It should be readily and completely soluble in Alcohol (90 p.c.).

1 c.c. of an aqueous solution of the salt when warmed with 1 c.c. of Potassium or Sodium Hydroxide Solution and sufficient Iodine Solution to colour the liquid distinctly brown yields a pale yellow precipitate of Iodoform.

1 c.c. of a saturated aqueous solution warmed with 2 c.c. of Potassium or Sodium Hydroxide and one or two drops of Aniline evolves the powerful, penetrating and highly poisonous odour of Phenyl isonitrile. It should volatilise completely when heated, and should leave no weighable residue upon ignition.

CHLORETONE ELIXIR—Chloretone, 10 grains, Spirit of Peppermint, 10 minims, Compound Tincture of Cardamoms, 1 fl. drim., Glycerin, q.s. to make 2 fl. drim., dilute immediately before use with 1 fl. oz. of Water for a dose.

Not Official

CHLORI LIQUOR

SOLUTION OF CHLORINE

Syn—AQUA CHLORI

A yellowish green liquid, possessing a powerful characteristic odour of Chlorine.

It is not now included in the text of the *B.P.*, but is transferred to the Appendix

Medicinal Properties—Deodouriser, antiseptic, and disinfectant. When diluted it is used as a gargle in smallpox, scarlatina, diphtheria, and putrid sore throat, and as a wash for ulcers, cancerous sores, buboes, and large abscesses.

Strongly advocated by Burney Yeo in the treatment of enteric fever. The solution he uses is obtained by pouring strong Hydrochloric Acid over Potassium Chlorate, thus into a 12 oz bottle put 30 grains powdered Potassium Chlorate, and pour on it 1 fl drim strong Hydrochloric Acid, cork, shake, and allow gas to generate, then add Water little by little till bottle is filled. He says it gives much better results and is more pleasant to take than the Liquor Chlori of the *B.P.* '85. To 12 fl oz of this solution he adds 24 to 36 grains of Quinine and 1 fl oz of Syrup of Orange peel, he gives 1 fl oz of this mixture every two, three, or four hours, according to the severity of the case. It is prescribed as **Mistura Chlori c Quinina** (*Burney Yeo*).

Further experiments on its use recorded in the Bradshaw Lecture on the treatment of enteric fever (*B.M.J.* '04, 11 1450) have shown that the tendency to intestinal fermentation is lessened, and the strength of the circulation well sustained, with corresponding benefit to the general aspect of the case.

Dose—10 to 20 minims = 0.6 to 1.2 cc, in a wineglassful of Water.

Incompatibles—Salts of Lead and Silver.

Antidotes.—In case of poisoning by Chlorine Water, the antidotes are White of Egg, Milk, Flour.

Foreign Preparation—Official in Austr (*Aqua Chlori*), not less than 0.4 pc; Russ and US (*Liquor Chlori Co*), about 0.4 pc; Ger (*Aqua Chlorata*), contains 0.4 to 0.5 pc of Gas, Dutch (*Solutio Chlori*), 0.4 to 0.425 pc; Swiss (*Chlorum Solutum*), 0.4 to 0.5 pc; Port (*Solutio de Chloro*), Span (*Agua de Cloro*) Not in Belg, Dan, Fr, Ital, Jap, Mex, Norw or Swed.

Tests—Solution of Chlorine should possess a specific gravity of 1.003, should first redden and then bleach blue Litmus paper, should immediately decolorise Indigo Sulphate Solution, should liberate Iodine from Potassium Iodide Solution. The latter reaction is utilised for the quantitative determination of the percentage of Chlorine. It should contain not less than 0.5 pc as determined by adding a measured quantity of 10 cc of the Liquor to a solution of 1 gramme of Potassium Iodide dissolved in 25 cc of Water, not less than 14.2 cc of Deci-normal Volumetric Sodium Thiosulphate Solution being required to decolorise the liquid, using Starch Solution as an indicator. It should leave no weighable residue upon evaporation.

LIQUOR CHLORI COMPOSITUS (U.S.)—Potassium Chlorate, granulated 5 grammes. Hydrochloric Acid, 18 cc, Distilled Water, to 1000 cc. Add 1 cc of Potassium Iodide solution and mix with 20 cc of Distilled Water to the Potassium Chlorate. The capacity of about 2000 cc. Insert in the flask a stopper perforated to admit a funnel of the capacity of about 100 cc containing a 10 grammes of purified Cotton well wetted with cold Water, place the flask on a water-bath containing boiling Water for a period of from 2 to 3 minutes, when the flask is completely filled with a greenish yellow gas remove it from the bath and add cold Distilled Water through the Cotton in the funnel in two separate portions of 500 cc each. After the addition of each separate portion of cold Distilled Water stopper the flask securely, invert, and thoroughly agitate the contents. This solution should be freshly made when wanted. Average dose—4 cc (1 fl drim).

LIQUOR CHLORI—Potassium Chlorate, 50 grains, Hydrochloric Acid, 100 minims, Water, to 1000 minims. Acid to the Chlorate in a large bottle, when the Chlorine given off is abundant, add Water gradually, corking and shaking the bottle. The solution is recommended (*B.M.J.* '93, 1 1004) for the preparation of 'Euchlorine' solution for use as a gargle in diphtheria, contains an excess of

Potassium Chlorate Place 20 to 30 grains of Potassium Chlorate in a dry 8 oz bottle with 10 minims of strong Hydrochloric Acid, the fumes will fill the bottle, which on the addition of Water with shaking will make a good solution

These ingredients in varying quantities are given in other Hospital Pharmacopœias

A Chlorine solution strongly recommended for irrigation of the fauces in diphtheria is made by pouring 5 minims of strong Hydrochloric Acid on 9 grains of powdered Potassium Chlorate, and gradually adding an oz of Water — *L '03, n 1774*

Sodium Chlorate is a tasteless salt, and answers equally well for the formation of 'Euchlorine'

GARGARISMA CHLORI —Potassium Chlorate, 200 grains, Strong Hydrochloric Acid, 40 minims, Water, to 20 fl oz Place the Potassium Chlorate in a dry bottle, pour the Acid upon it, and set aside, loosely corked, for 10 minutes Then add the Water in 4 or 5 successive portions, shaking between each addition, so that the gas may be absorbed as completely as possible

Note —This gargle is usually employed diluted with one or more parts of Water It should be recently prepared as it deteriorates slowly on standing, and quickly if exposed to light —*St Thomas's*

The quantities given in the *B P C* are —Potassium Chlorate, 2 25, Hydrochloric Acid, 0 50, Distilled Water, sufficient to make 100 00 Same directions as above

VAPOR CHLORI (*B P* 1885) —Chlorinated Lime, 2 oz, cold Water, a sufficiency Put the powder into a suitable apparatus, moisten it with the Water and let the vapour that arises be inhaled

CHLOROFORMUM.

CHLOROFORM

TRICHLORO METHANE, METHENYL TRICHLORIDE

CHCl₃, eq 118 48

FR, CHLOROFORME, GER, CHLOROFORM, ITAL, CLOROFORMIO, SPAN.,
CLOROFORMO

A clear, colourless, heavy, mobile liquid, possessing a distinctive ethereal odour, and a sweet burning taste

It may be prepared by the action of Chlorinated Lime on Ethyl Alcohol or on Acetone The product obtained by its action on Methylated Alcohol is known as Methylated Chloroform

The *U S P* defines Chloroform as a liquid consisting of from 99 to 99 4 p c by weight of absolute Chloroform, and 0 6 to 1 0 p c of Alcohol, but the quantity of Alcohol is not now defined in the *B P* except that the product is worked to a specific gravity 1 490 to 1 495

It should be kept in well stoppered amber tinted glass bottles, in a cool place, and protected as far as possible from the light

Solubility —10 in 7 of Alcohol (90 p c), in all proportions of Ether and Alcohol, freely in Olive Oil and Oil of Turpentine In Water at 32° F 1 in 150, at 60° F 1 in 185, at 86° F 1 in 210, at 113° F 1 in 200, at 130° F 1 in 192 Will not dissolve in Glycerin

Chloroform acts on Vulcanite, and dissolves Caoutchouc, Gutta-percha, Mastic, Elemi, Tolu, Benzoin and Copal Amber, Sandarach, Lac and Beeswax are only partially soluble It also dissolves Iodine, Bromine, most of the alkalis, the fixed and volatile Oils, most Resins and Fats It dissolves Sulphur and Phosphorus sparingly

Medicinal Properties. A general anæsthetic Internally a sedative, carminative and antispasmodic Its chief use is to produce general anæsthesia by **inhalation** during surgical operations, uræmic and puerperal convulsions, and in obstetric practice Should be given with great caution in cases of fatty and dilated heart, in extensive lung disease and severe anæmia Internally, useful to relieve flatulent distension of stomach and bowels, and the cough of fibroid phthisis, in delirium tremens, and in sea-sickness Externally, with Camphor, relieves toothache and neuralgia Applied immediately after the sting of a wasp, takes away the pain A powerful auxiliary to the Liniments of Aconite and Belladonna

Its vapour and aqueous solution are antiseptic, and the addition of 1 minim to 1 fl oz of animal or vegetable infusion will preserve it Vinegar after Chloroform inhalation to prevent sickness (See p 9)

Chloroform should not be used as an anæsthetic in a room where gas is being burned, a mixture of Chloroform vapour and air being decomposed by a flame with the formation of irritating compounds—*L* '99, i 1728, *T* (1909), 601, *P J* '02, i 876

The dosage of Chloroform for inhalation A powerful and dangerous anæsthetic, not to be recommended in minor surgery The notorious uncertainty and danger in Chloroform administration is the uncertainty in the quantity administered 0.2 c.c. per minute recommended, in a mixture of Chloroform and air at an average percentage of 1.5 p.c.—*B M J* '98, i 1057-1062

An attempt to ensure its admixture with air in certain proportions is described in *L* '04, ii 1462 The Report of the Chloroform Committee was presented at the Oxford meeting of the British Medical Association, the Vernon Harcourt inhaler was introduced, and the adoption of a maximum strength of 2 p.c. Chloroform vapour as an adequate and safe limit for general surgical purposes recommended (*B M J* '08, ii 141, '04, ii 161, *L* '04, ii 1856), a further reference to the subject is made (*J C S Abs* 1904, ii 756), showing all that was necessary in the air was 2 p.c., and that danger lay beyond It has latterly been shown (*J C S* 1904, *Trans* 949, *L* '05, i 589) that Chloroform derived from Acetone is inferior in anæsthetic properties to Chloroform derived from Alcohol The difference is stated to be due to the presence of 0.05 p.c. Ethyl Chloride Acetone Chloroform to which this amount of Ethyl Chloride had been added, exhibited equally as good anæsthetic effects as Chloroform prepared from Ethyl Alcohol This statement has, however, been challenged by Messrs J F MacFarlan & Co (*L* '05, i 747), who think that Chloroform prepared from Acetone is by no means generally accepted as inferior to Chloroform prepared from Alcohol, and that up to the present insufficient evidence has been adduced to establish these views

Less Chloroform is required when preceded by Morphine Scopalamine injections (See p 650)

Dose—1 to 5 minims = 0.06 to 0.3 c.c.

Ph. Ger. maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes

Prescribing Notes—Chloroform Water and Spirit of Chloroform are used as sweetening agents, and to preserve solutions from decomposition As a rule in 'mixtures' Chloroform is in such small quantities as to dissolve in the Water, in concentrated 'mixtures' Mucilage of Gum Acacia would be required to suspend it, it can be given in 'drops' dissolved in some strongly alcoholic menstruum

It is used with Camphor Liniment, Soap Liniment, Olive Oil, or Oil of Peppermint

Camphorated Chloroform, see Camphor

Official Preparations—Aqua Chloroformi, Linimentum Chloroformi, Spiritus Chloroformi Tinctura Chloroformi et Morphinæ et Scopolaminæ

Not Official—Chloroethoform, Chlorodyspnum, Fumario Chloroformi, Liquor

Chloroformi Compositus, Mistura Tussi Rubra Concentrata, Mistura Chloroformi Composita, Paroenum Chloroformi Camphoratum, Pomade de Chloroforme, Tinctura Chloroformi Composita, Chloroformum Camphoratum, Carbon Tetra chloride, A C E Mixture, Vienna Mixture, 'Methylene,' Regnaud's Anæsthetic Mixture, Pental, Vapor Chloroformi Compositus, Vasolimentum Chloroformi Camphoratum

Antidotes—In case of overdose of Chloroform, the antidotes are, fresh pure air and artificial respiration (*M T* '74, ii 219), and Amyl Nitrite—*L* '75, i 644 *B M J* '97, ii 352 Hypodermic injection of Starchine, altogether $\frac{1}{2}$ grain was used in divided doses of $\frac{1}{2}$ grain followed by $\frac{1}{2}$ grain—*B M J* '97, ii 1498

Foreign Pharmacopœias—Official in Austri and Belg, sp gr 1 485 to 1 490, Fr sp gr 1 495 to 1 500, U S, sp gr not below 1 476 at 25° C (77° F), Dan, Dutch, Ger, Hung, Norw, Swed and Swiss, sp gr 1 485 to 1 489, Ital, sp gr 1 490 to 1 493, Jap, sp gr 1 485 to 1 495, Mex, Port and Span, sp gr 1 480, Russ, sp gr 1 499 to 1 500

The new Austri, Dutch and Swiss Pharmacopœias include Chloroform and Chloroform pro narcosi. The Austri and Swiss Chloroform and Chloroform pro narcosi are required both to possess the same sp gr, the Dutch Chloroform possesses the sp gr 1 485 to 1 489, that 'pro narcosi' a sp gr of 1 498 to 1 500, and is prepared by the decomposition of Chloral Hydrate by Sodium Hydroxide. Fr has Chloroforme rectifié du commerce sp gr 1 495 to 1 500 which must not be used as an anæsthetic, also Chloroforme anæsthésique sp gr 1 498 which is prepared from the other

Tests—The distinguishing tests for Chloroform are the specific gravity, which is about 1 490, and the boiling point, which should be about 61° C (141 8° F). Pure Chloroform has a specific gravity of 1 5, and boils at 60 8° C (141 44° F). The *BP* gives the specific gravity as 1 490 to 1 495, the *USP* not below 1 476 at 25° C (77° F), the *PG* 1 485 to 1 489. The *BP* and the *PG* give a similar boiling point 60° to 62° C (140° to 143 6° F), the *USP* gives 60° to 61° C (140° to 141 8° F). When boiled with Potassium or Sodium Hydroxide Solution it is decomposed yielding a solution which gives with Silver Nitrate Solution, when acidified with Nitric Acid, a white curdy precipitate soluble in Ammonia Solution and which is rapidly blackened on heating. A few drops of Chloroform when warmed with 2 or 3 cc of an alcoholic Potassium or Sodium Hydroxide Solution and a drop or two of Aniline evolve the powerful, penetrating and highly poisonous odour of Phenyl-isocyanide. A few drops of Chloroform, when heated with Fehling's (Potassio-cupric Tartrate) Solution, yield a reddish deposit of Cuprous Oxide.

The more generally occurring impurities are pyrogenous oils, acid, free Chlorine, Chlorides, secondary products of decomposition and fixed matter. Acid, free Chlorine, and Chlorides are all extracted by means of Water, and if the sample be shaken for 5 minutes with twice its volume of Water, free Acid may be detected in the aqueous liquid by its reaction towards blue Litmus paper, Free Chlorine by any blue coloration produced on the addition of 1 cc of Cadmium Iodide Solution and 2 drops of Starch Mucilage, and Chlorides by the addition of Silver Nitrate Solution, when not more than a very slight opalescence should be produced. In testing for Free Chlorine the *BP* uses Cadmium Iodide Solution, the *USP* Potassium Iodide Test Solution, and the *PG* Zinc Iodide Solution. Pyrogenous oils may be detected by allowing a definite volume to evaporate from a large piece of filter paper placed on a warm plate, when no foreign odour

should be perceptible. The *BP* employs 20 c.c. for this test, the *USP* 10 c.c., whilst the *PG* specifies no particular quantity. The *BP* requires that no odour should be perceptible at any stage of the evaporation, the *USP* that an odour should not be perceptible during the later stages of the evaporation and the filter paper should remain odourless, the *PG* that the filter paper shall not retain any foreign odour after the evaporation of the Chloroform. Secondary products of decomposition may be detected by the behaviour of the specimen with concentrated Sulphuric Acid. After shaking the acid with 10 times its volume of Chloroform for 20 minutes, and allowing the mixture to remain at rest for 15 minutes, both the acid and the chloroformic layers should be perfectly transparent and nearly colourless, a portion of the Sulphuric Acid layer diluted with Water should remain transparent, should be very nearly colourless and should possess no disagreeable odour, and if the liquid be further diluted and tested with Silver Nitrate Solution no more than a diminished transparency should be caused. The *USP* divides the test into three separate headings indicating impurities decomposable by Sulphuric Acid, odorous decomposition products, and chlorinated decomposition compounds. Fixed matter may be detected by any residue remaining after the evaporation of the Chloroform.

Starch Solution and Cadmium, Potassium or Zinc Iodide—The aqueous portion obtained by shaking Chloroform with twice its volume of Water for 5 minutes, should not afford any colour with 1 c.c. of Cadmium Iodide T.S. and 2 drops of Mucilage of Starch, *BP*, when Chloroform is shaken with Zinc Iodide and Starch Solution, the Starch Solution should not become blue, nor the Chloroform coloured, *PG*. The *USP* requires that the aqueous solution should not become coloured with T.S. of Potassium Iodide.

Silver Nitrate—The addition of 4 drops of Silver Nitrate Solution should not produce more than a very slight opalescence in the aqueous portion obtained by shaking Chloroform with twice its volume of Water for 5 minutes, *BP*, should not be affected by T.S. of Silver Nitrate, *USP*. It should not yield any turbidity with T.S. of Silver Nitrate diluted with as much Water, *PG*.

Sulphuric Acid—Chloroform when shaken with one-tenth of its volume of Sulphuric Acid for 20 minutes and set aside for 15 minutes should acquire practically no colour in either the chloroformic or Sulphuric Acid layer, and both should be quite transparent, *BP*, the *USP* directs the use of 4 c.c. of Sulphuric Acid and 40 c.c. of Chloroform shaken in a 50 c.c. glass stoppered cylinder during 5 minutes, and that the liquids be allowed to separate completely so that both are transparent. The Chloroform should remain colourless and the acid should appear colourless or very nearly so when in a stratum of not less than 15 mm. in thickness. In the *PG* test 15 c.c. of acid are shaken with 20 c.c. of Chloroform in a stoppered glass cylinder of 3 cm. diameter previously rinsed out with Sulphuric Acid. The acid should not become coloured within 1 hour.

2 c.c. of the Sulphuric Acid liquid obtained on shaking the Chloroform as above described, when diluted $2\frac{1}{2}$ times its volume of Water should remain clear, and should possess a pleasant odour, *BP*, should be colourless and clear, and hot from the mixing should be odourless or give but a faint vinous or ethereal odour, *USP*. The liquid obtained in carrying out the *BP* test, immediately above, should still retain its transparency and freedom from colour, even when further diluted with 10 c.c. of Water and stirred with a glass rod and the transparency should not be more than slightly diminished on the addition of 4 drops of Silver Nitrate Solution, *BP*, should remain clear and

should not be affected by T S of Silver Nitrate, *U S P*. The addition of Silver Nitrate Solution should not more than slightly diminish the transparency of the aqueous portion obtained by shaking the Sulphuric Acid treated Chloroform with twice its volume of Water, *B P*.

Volumetric Determination—Chloroform may be determined in the absence of other reducing substances by Fehling's Solution. A more accurate method is to pass the vapour through a red hot tube containing Platinum wire gauze, which decomposes the Chloroform with the formation of Hydrochloric Acid. The products are collected in a bulb tube containing Water, and the acid produced is titrated with Volumetric Potassium or Sodium Hydroxide Solution. 108.57 parts by weight of Hydrochloric Acid are equivalent to 118.48 parts by weight of Chloroform. This method has been applied to the determination of Chloroform in animal tissues (*B M J* '01, 11 1859), but the Hydrochloric Acid is recommended to be determined with Volumetric Silver Nitrate Solution, the excess of Silver being determined with Volumetric Ammonium Thiocyanate Solution, using Ferrous Ammonium Sulphate Solution as an indicator. In using the combustion method of determination for Chloroform in blood, the blood should be mixed with an equal volume of a saturated aqueous solution of Urea, by which means (*B M J* '03, 11 cxlii) the blood remains fluid during the necessary heating and more than 90 per cent of the Chloroform is accounted for.

Preparations

AQUA CHLOROFORMI CHLOROFORM WATER

Chloroform, 30 minims, Distilled Water, *qs* to make 25 fl oz
(1 in 400)

Half the strength of *B P* 1885

Dose — $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc, but ordered in smaller quantities as a flavouring agent

Foreign Pharmacopœias—Official in Austr (1 in 100), Belg, Dan, Fr, Norw and Swiss (1 in 200), Dutch, Solutio Chloroformi Aquosa (1 in 250), Ital (1 in 2000), Jap (1 in 400), Span (1 in 250), U S, Saturated Aqueous Solution. Not in the others.

Tests—Chloroform Water has a specific gravity of 1.001, should be neutral in reaction to Litmus paper, and when warmed with a little Alcoholic Potassium or Sodium Hydroxide Solution and a few drops of Aniline should evolve the powerful, penetrating, and highly-poisonous vapours of Phenyl-isocyanide.

LINIMENTUM CHLOROFORMI LINIMENT OF CHLOROFORM.

Chloroform, 1, Liniment of Camphor, 1 (1 in 2)

The Oil in the Camphor Liniment prevents rapid evaporation of the Chloroform.

Foreign Pharmacopœias—Official in Ger and Jap (*Oleum Chloroformi*), Chloroform 1, Olive Oil 1, Austr (*Linimentum Chloroformiatum*), Chloroform, Oleoso balsamic Mixture, Spirit of Ether, Spirit of Camphor, Spirit of Potash Soap, equal parts of each, Fr (*Liniment au Chloroforme*), Chloroform 1, Poppy Oil 9, Swed (*Linimentum Chloroformi Comp*), Chloroform 3, Camphor 3, Alcohol (90 pc) 5, Camphorated Soap Liniment 6, Tincture of Opium 3, Swiss (*Oleum Chloroformi*), Chloroform 1, Olive Oil 3. Jap has also *Linimentum Chloroformi*, Chloroform 1, Camphor Oil 1. All by weight. U S, Chloroform 3, Soap Liniment 7. Not in the others.

Tests—It should possess a specific gravity of 1.212, and contains about 37 pc w/v of total solid residue.

SPIRITUS CHLOROFORMI SPIRIT OF CHLOROFORM *B P Syn*
 —CHLORIC ETHER, SPIRIT OF CHLORIC ETHER

Chloroform, 1, Alcohol (90 p c), *q s* to make 20 (1 in 20)

Dose.—5 to 20 minims = 0.3 to 1.2 c c, for repeated administration, for a single administration 30 to 40 minims = 1.8 to 2.4 c c

Frequently prescribed as a sweetening agent, and to cover nauseous flavour.

Foreign Pharmacopœias —Official in Jap, 1 in 20, U S, Chloroform 6, Alcohol 94 Not in the others

Tests —Spirit of Chloroform has a specific gravity of about 0.860, and should leave no weighable residue on evaporation

TINCTURA CHLOROFORMI ET MORPHINÆ COMPOSITA.
 COMPOUND TINCTURE OF CHLOROFORM AND MORPHINE

Chloroform, 1½ fl oz, Morphine Hydrochloride, 87½ grains, Diluted Hydrocyanic Acid, 1 fl oz, Tincture of Capsicum, ½ fl oz, Tincture of Indian Hemp, 2 fl oz, Oil of Peppermint, 14 minims, Glycerin, 5 fl oz, Alcohol (90 p c), *q s* to make 20 fl oz (about 1 in 13)

Dose.—5 to 15 minims = 0.3 to 0.9 c c

10 minims contain Chloroform ¾ minim, Morphine Hydrochloride, 1½ grain, Diluted Hydrocyanic Acid ½ minim, Tincture of Indian Hemp 1 minim

It is nearly 4½ times stronger in Morphine than *B P* 1885, and in other respects differs considerably

The *B P* 1885 preparation was practically the same as *Liquor Chloroformi Compositus*, except that the former contained four times as much Morphine as

In *B P* 1898 the formula was completely changed, therefore *Liquor Chloroformi Compositus*, previously omitted in *Companion*, is re-inserted

The *B P C* have also added a formula under the name 'Chlorodyne' (see below)

Foreign Pharmacopœias —Official in Jap. Hung has a 'Chlorodyne', but it differs considerably from *B P* Not in the others

Tests.—Compound Tincture of Chloroform and Morphine has a specific gravity of 1.010 to 1.015, it should contain from 28.0 to 30.0 p c w/v of total solids and from 52 to 54 p c w/v of Absolute Alcohol

Not Official

LIQUOR CHLOROFORMI COMPOSITUS (*Squire*) —Chloroform, 4 fl oz, Ether, 1 fl oz, Alcohol (90 p c), 4 fl oz, Treacle, 4 fl oz, Extract of Liquorice, 2½ oz, Morphine Hydrochloride, 8 grains, Oil of Peppermint, 16 minims, Syrup, 17½ fl oz, Prussic Acid (2 p c), 2 fl oz Mix the Oil of Peppermint, Alcohol and Prussic Acid together, and dissolve the Morphine Hydrochloride in the mixture, add the Chloroform and Ether, dissolve the Extract of Liquorice in the Syrup, add the Treacle, and mix in the other ingredients

This formula first appeared in the *Companion* in 1864

Dose —5 to 10 minims = 0.3 to 0.6 c c

10 minims contain Chloroform about 1 minim, Diluted Hydrocyanic Acid ½ minim, Morphine Hydrochloride 1½ grain

CHLORODYNUM (*B P C*) —Chloroform, 6.00 Morphine Hydrochloride, 0.50, Tincture of Indian Hemp, 8.00, Tincture of Capsicum, 1.50, Liquid

Extract of Liquorice, 12 00, Mucilage of Acacia, 12 00, Tincture, 20 00, Glycerin, 22 00, Oil of Peppermint, 0 10, Alcohol, sufficient to produce 100 00

10 minims contain Chloroform about $\frac{1}{2}$ minim, and Morphine Hydrochloride about $\frac{1}{10}$ grain, and no Hydrocyanic Acid

The *B P C* Chlorodyne Lozenges contain about $\frac{1}{15}$ grain of Morphine Hydrochloride

EMULSIO CHLOROFORMI—Chloroform, 1 fl oz, Tincture of Quillaia, 1 fl drim, Water, to 20 fl oz—*London*

Chloroform, 1 fl oz, Tincture of Quillaia, 3 fl drim, Water, to 20 fl oz—*St Thomas*

The *B P C* quantities are very similar Chloroform, 5, Tincture of Quillaia, 2, Distilled Water, to make 100

MISTURA TUSSI RUBRA CONCENTRATA—Diluted Hydrobromic Acid, 15 minims, Compound Tincture of Chloroform, 10 minims, Compound Tincture of Cardamoms, 10 minims, Solution of Morphine Hydrochloride, 5 minims, Diluted Hydrocyanic Acid, 1 minim, Syrup of Wild Cherry, to 1 fl drim—*Australian Pharmaceutical Formulary*

Mistura Chloroformi Composita *Syn* Mistura Tussi Rubra—Morphine Hydrochloride, $\frac{1}{8}$ grain Diluted Hydrobromic Acid, 30 minims, Chloroform, $\frac{1}{2}$ minim, Tincture of Cudbear, 7 $\frac{1}{2}$ minims, Syrup of Wild Cherry, 90 minims, Syrup, to 2 fl drim—*B P C*

This formula has been changed very considerably in the *B P C Supplement*, and as now amended reads—Morphine Hydrochloride, 0 05, Diluted Hydrobromic Acid, 12 5, Cherry laurel Water, 3, Chloroform, 0 25, Syrup of Tolu, 25, Tincture of Cudbear, 9, and sufficient Syrup to produce 100

POMMADE DE CHLOROFORME—Chloroforme rectifié 10, Yellow Wax 5, Lard 85—*F*

TINCTURA CHLOROFORMI COMPOSITA—Chloroform, 2, Alcohol (90 p c), 8, Compound Tincture of Cardamoms, 10 (1 in 10)

It first appeared in *B P* 1885, but was omitted in 1898, and subsequently included in *B P C*

Dose—5 to 60 minims = 0 3 to 3 6 c c

The Chloroform will separate if this Tincture is prescribed in too little Water Has been given successfully for the prevention of sea-sickness

CHLOROFORMUM CAMPHORATUM—Camphor, 2, Chloroform, 1, dissolve—*B P C Formulary* 1901, incorporated in *B P C*

A remedy for toothache, and topically applied for rheumatism

ACE MIXTURE—Alcohol (90 p c), 1, Chloroform, 2, Ether, 3, mix

Used as an anæsthetic in place of Chloroform—*Med Chir Trans* vol 47, '64, 341, *B M J* '87, ii 975, 1078, 1185, 1314, 1359 Advantages over Chloroform—*B M J* '97, ii 160

ACE (*Martindale*)—Absolute Alcohol (sp gr 0 795), 1, Chloroform (sp gr 1 497), 2, Purified Ether (sp gr 0 720), 3

VAPOR CHLOROFORMI COMPOSITUS (*B P C*)—Alcohol (90 p c), 4, Chloroform, 8, Purified Ether, *q s* to make 25

VASOLIMENTUM CHLOROFORMI CAMPHORATUM—Camphor, 3, Chloroform, 3, Liquid Vasoliment (*see* p 864), 3, all by weight—*Hager*

Parogenum Chloroformi Camphoratum *Syn* Camphorated Chloroform Vasoliment—Camphor, 3, Chloroform, 2, Parogen (*see* p 864), 3—*B P C*

VIENNA MIXTURE—Ether, 3, Chloroform, 1, by weight

'METHYLENE' (formerly called Methylene Bichloride)—Introduced by B W Richardson in November, 1867 It is a limpid, dense fluid, sp gr varies, when dropped into Water about one fourth of it is dissolved, the remainder separates like Chloroform at the bottom of the vessel as a perfectly clear and

distinct fluid, and the whole has a sweet, pleasant odour, without the least smell of Ether

Recommended as an anæsthetic in place of Chloroform

REGNAULD'S ANÆSTHETIC MIXTURE—Chloroform, 4, Methyllic Alcohol, 1, mix

Used as an anæsthetic in place of Chloroform

CHLORÆTHOFORM—Chloroform (from Acetone), 100, Ethyl Chloride, 0.25

CARBON TETRACHLORIDE—A colourless, volatile, heavy liquid. It may be prepared by the action of dry Chlorine gas on Carbon Bisulphide vapour, or by replacing the Hydrogen ion of Chloroform with the Chlorine ion

It should be kept in dark amber-tinted glass-stoppered bottles, and in a cool atmosphere

Has been employed to produce anæsthesia, but its principal use is as an inhalation in hay fever, and as an application on Pline for neuralgia

Danger attending its use as a hair-wash (*B M J* '07, ii 764, 776), or dry shampoo—*L* '07, i 1709

Tests—Carbon Tetrachloride has a specific gravity of 1.599 to 1.600 and a boiling point of 77° to 78° C (170.6° to 172.4° F). When warmed with Potassium or Sodium Hydroxide Solution, it is decomposed with the formation of Potassium or Sodium Chloride and Carbonate, on the addition of dilute Nitric Acid, and the faintly acidified Silver Nitrate Solution a white precipitate, insoluble in Nitric Acid, but soluble in Ammonia Solution. When warmed with an alcoholic Potassium or Sodium Hydroxide Solution and a few drops of Aniline it evolves the powerful, penetrating, and highly-poisonous vapours of Phenol-isocyanide

It may be contaminated with other Chlorine compounds or with chlorinated decomposition products, *e.g.*, Hydrochloric Acid. The former may be detected by the darkening in colour produced when a portion of the specimen is mixed with an equal volume of concentrated Sulphuric Acid, the latter by shaking the sample with twice its volume of Water and noting the reaction of the aqueous liquid towards Litmus paper and Silver Nitrate Solution, it should neither redden blue Litmus paper nor should it produce a turbidity with Silver Nitrate Solution

PENTAL (Trimethylethylene)—A colourless, mobile, inflammable liquid. Has been recommended as a general anæsthetic for short operations. Whittle states that several deaths have been attributed to it, and that it causes albuminuria—*M A* '95, 40, *L* '94, i 1080, '96, i 45, 710, 950, *T G* '93, 34, '94, 555, *B M J E* '93, ii 28, *B M J* '96, i 730

CHRYSAROBINUM.

CHRYSAROBIN

FR, CHRYSAROBINE, GER, CHRYSAROBINUM, ITAL, CRISAROBINA, SPAN, CHRYSAROBINA

An odourless and tasteless, yellow, crystalline powder, obtained from Araoba

Purified Chrysarobin was introduced into medicine incorrectly as **Chrysophanic Acid**, and it is still known by this name, which, however, only correctly applies to the oxidised product

Araoba yields from 55 to 80 p.c. (average 71 p.c.) of Chrysarobin—*P J* (3) xxii 544

Medicinal Properties.—In form of **unguentum** or **pigmentum**, it has been found efficient in chronic psoriasis, and is a powerful parasiticide in ringworm and other parasitic skin diseases, but as it may cause erythema it requires watching, it should not be allowed to touch the healthy skin. The ointment stains the skin

yellow, also the linen Has been given internally for psoriasis, eczema and acne, but it is very irritating, producing purging, griping and vomiting even in very small doses

To remove the stains from linen first remove all grease with Benzol and then apply a solution of Chlorinated Lime In some cases a little Caustic Soda solution also may be necessary—*C D* '99, 1 652

Alopecia areata, treated almost exclusively with Chrysarobin stick—Chrysarobin, 80, Colophony Resin, 5, Yellow Wax, 35, Olive Oil (by weight), 80 (*B M J E* '95, 11 103), and with excellent result by ointment 2 dm to oz—*B M J* '07, 11 491

Chrysophanic Acid is not an efficient substitute for Chrysarobin in the treatment of psoriasis—*B M J E* '96, 11 96

Used in the form of an ointment (*B M J* '05, 1 699) either alone or combined, according to circumstances, with Tar or Salicylic Acid, it cures chronic psoriasis

Dose $\frac{1}{10}$ to 1 grain = 0.006 to 0.06 gramme

Official Preparation—Unguentum Chrysarobini

Not Official—Unguentum Chrysarobini Compositum, Pigmentum Chrysarobini, Chrysarobin Plaster Mulls, Anthrarobin, Eurobin, Lenirobin

Foreign Pharmacopœias—Official in Austr (Araroba Depurata), Belg, Dan, Dutch, Ger, Ital, Jap, Norw, Russ, Swed, Swiss and US (Chrysarobinum), Mex (Crisarobina), the purified product Not in the others

Tests—Chrysarobin melts, according to the *USP*, at about 157° C (314.6° F) It varies in its behaviour towards different solvents, in Water it is only slightly soluble, in Petroleum Spirit it partly dissolves It is almost completely soluble in hot Alcohol (90 p c) and completely soluble in Chloroform, the *BP* mentions that it partially dissolves in Potassium Hydroxide Solution, assuming a deep brownish-red colour, the *USP* that it is soluble in dilute or concentrated Potassium Hydroxide solutions, forming a red coloured liquid with green fluorescence, the *PG* states that when boiled with 2000 parts Water it does not completely dissolve and yields a filtrate which does not affect Litmus paper, and which is not affected by Ferric Chloride Test-solution It dissolves to a reddish-yellow solution in concentrated Sulphuric Acid and is reprecipitated unchanged on dilution with Water A carmine red colour should be assumed in the course of a day by Ammonia Solution which has been shaken with Chrysarobin It may be distinguished from Chrysophanic Acid by mixing 1 mgm with 2 drops of fuming Nitric Acid and adding Ammonia Solution, a violet coloration is produced, whereas Chrysophanic Acid produces a yellow coloured liquid, when Chrysarobin is shaken with Lime Water for a few minutes a violet coloration is imparted to the liquid When heated in an open crucible it melts, and when ignited with free access of air it is officially required not to leave more than 1 p c of ash, the *PG* says 0.2 gramme should leave no weighable residue, the *USP* says it is entirely consumed

Preparation

UNGUENTUM CHRYSAROBINI. CHRYSAROBIN OINTMENT

2 of Chrysarobin dissolved in 48 of Benzoated Lard by the aid of heat, and subsequently stirred till cold (1 in 25)

Official in US, 1 in about 17

Not Official

UNGUENTUM CHRYSAROBINI COMPOSITUM (*Unna*)—Chrysarobin and Ichthyol, of each, 5, Salicylic Acid, 2, Yellow Vaseline, 88

PIGMENTUM CHRYSAROBINI—Chrysarobin, 1, Gutta Percha solution, 9—*Guy's*

Chrysarobin, 60 grains, Chloroform, 10 dm, pure Gutta Percha, 60 grains, dissolve Painted on with a stiff brush Acts effectually, and does not stain the linen—*B M J* '87, 11 1139

A 5 to 10 p c solution of Chrysarobin in equal parts of Chloroform and Glycerin Used in ringworm, applied till erythema and a slight oedema are produced—*B M J* '04, 1 16

Chrysarobin, 1, solution of Gutta Percha, *q s* to make 10—*B P C*

CHRYSAROBIN PLASTER MULLS (*Unna*)—Contain $\frac{1}{10}$ grain to the square inch, also five times this strength

ANTHRAROBIN ($C_{14}H_{10}O_2$, eq 224.38)—A yellow, or light yellowish brown, odourless, tasteless powder. A reduction product from Alizarin. Slightly soluble in Water, but readily in Alcohol (90 p c) and solution of Borax

Tests—The aqueous solution yields with Lead Acetate solution a reddish brown precipitate, with Ferric Chloride Solution, a brownish-violet precipitate. It dissolves in Sodium Hydroxide Solution (15 p c) with the production of a brownish-yellow colour which changes to violet on absorption of Oxygen from the air. It should leave not more than 2 p c of residue on incineration.

A substitute for Chrysarobin. For an ointment it is rubbed with Olive Oil and diluted with Lard.

Its action is similar to Chrysarobin, but it is slower and does not produce the same irritation. The part should be previously washed with Potash Soap, and the alcoholic tincture is preferred to the ointment. The strength of the ointment used is 1 in 10—*B M J* '88, 1 1234, *L M R* '88, 234, and '89, 243

Eurobin and **Lenirobin** are Chrysarobin Acetates, soluble in Acetone and Chloroform, they have been used for the same purposes as Chrysarobin. It is stated that they do not stain the normal skin, or the linen, like Chrysarobin.

CIMICIFUGÆ RHIZOMA.

CIMICIFUGA

B P Syn—ACTÉE RACEMOSÆ RADIX

FR, RACINE D'ACTÉE À GRAPPES, GER, SCHLANGENWURZEL

The dried Rhizome and Roots of *Cimicifuga racemosa*

The active principle is probably a resinous amorphous substance

Medicinal Properties—Bitter stomachic, analgesic. Given in neuralgia, myositis, rheumatism, " " " " and sciatica. Relieves the pain of dysmenorrhœa and pleurodynia.

Official Preparations—Extractum Cimicifugæ Liquidum, and Tinctura Cimicifugæ

Not Official—Cimicifugin

Foreign Pharmacopœias.—Official in U S. Not in the others.

Descriptive Notes.—The size of the official rhizome of *Cimicifuga racemosa*, Ell, is $\frac{1}{2}$ to 1 inch (12 to 25 mm) in diameter and 2 to 6 inch (5 to 15 cm) long, with slightly curved branches marked with transverse leaf scars, and the remains of ascending stems. In transverse section the large horny pith is

surrounded by a zone of narrow woody wedges and large medullary rays, and a relatively thin bark. The rootlets, which are usually more or less broken off, show in transverse fracture about four woody wedges arranged like a Maltese cross, set in a dark cortical portion. The taste is bitter and acid. It is sometimes confused with *Helleborus niger*, L., but that rhizome is smaller, $\frac{1}{10}$ to $\frac{1}{10}$ inch (5 to 7.5 mm), and has more erect branches, with short woody wedges 8 to 12 in number and a thick bark, and the woody wedges in the roots taper outwards so that the central column has a cylindrical and stellate appearance.

Tests—*Cimicifuga* may be distinguished from Black Hellebore by Ferric Chloride Test-solution, the *BP* says that the Rhizome and Roots are blackened by the reagent, but the colour is really a greenish black. No official limit of ash is given, but it generally leaves about 5 to 7 p.c. of ash and it should not amount to more than 10 p.c.

Preparations

EXTRACTUM CIMICIFUGÆ LIQUIDUM LIQUID EXTRACT OF CIMICIFUGA *BP Syn*—LIQUID EXTRACT OF *ACTÆA RACEMOSA*

20 of *Cimicifuga* percolated with Alcohol (90 p.c.) until exhausted, reserving the first 15, and evaporation of the further portion to a soft extract which is dissolved in the 15, and the whole made up to 20 with Alcohol (90 p.c.) (1 in 1)

Dose—5 to 30 minims = 0.3 to 1.8 c.c.

Foreign Pharmacopœias—Official in U.S., also a powdered extract prepared by evaporation of the fluid extract and admixture with powdered Licorice root. Not in the others.

Tests—Liquid Extract of *Cimicifuga* has a specific gravity of 0.875 to 0.890, it contains from 8 to 10 p.c. w/v of total solids and about 80 p.c. w/v of Absolute Alcohol.

TINCTURA CIMICIFUGÆ TINCTURE OF CIMICIFUGA *BP Syn*—TINCTURE OF *ACTÆA RACEMOSA*

2 of *Cimicifuga*, in No. 40 powder, percolated with Alcohol (60 p.c.), to yield 20 (1 in 10)

Dose—30 to 60 minims = 1.8 to 3.6 c.c.

The Tincture formerly in the *Companion* as 'Not Official' was twice the strength of this, and is still ordered as *Tinctura Actææ Racemosæ (Square)* to distinguish it from the Official Preparation.

Foreign Pharmacopœias—Official in U.S., 1 in 5. Not in the others.

Tests—The specific gravity of the tincture should be between 0.918 to 0.925, it contains from 1.0 to 2.8 p.c. w/v of total solids and about 58 p.c. w/v of Absolute Alcohol.

Not Official

CIMICIFUGIN—A brown powder, a large proportion of which is soluble in Alcohol (90 p.c.)

Dose—1 to 5 grains = 0.06 to 0.32 gramme

It is stated in *USD* that it is an impure Resin obtained by precipitating a saturated tincture of the root with Water.

Not Official

CINCHONÆ CORTEX

CINCHONA BARK

The dried Bark of *Cinchona Calisaya*, *C. officinalis*, *C. lancifolia*, and other species of *Cinchona*, from which the various alkaloids of the bark may be obtained

The official salts of Quinine, which are Quininæ Hydrochloridum, Quininæ Hydrochloridum Acidum, and Quininæ Sulphas, may be prepared from the Bark of various species of *Cinchona* and *Remijia*

Only Red *Cinchona* Bark is official for the galenical preparations

Foreign Pharmacopœias—Official in Austr., Dan., Jap., Norw., Russ. and Swed., any species, especially *Succirubra*, Dutch, Ger. and Swiss (*Cinchona Succirubra*), Fr. (*Quinquina Jaune* and *Quinquina Rouge*), Mex., any species, Hung. (*China Calisaya* and *Succirubra*), Port. (*Cinchona Flava*, *Fusca* and *Rubra*), Span. (*Cinchona Calisaya*, *Peruviana* and *Succirubra*), Belg., Swiss and Ital. (*Cinchona Succirubra*, *Ledgeriana* and *Calisaya*), U.S., any species of *Cinchona*, especially *Ledgeriana*, *Calisaya* and *Officinalis*, the latter used for Compound Tincture only

CINCHONÆ RUBRÆ CORTEX.

RED CINCHONA BARK

FR., QUINQUINA ROUGE, GFR., CHINARINDE, ITAL., CHINA ROSSA,
SPAN., QUINA ROJA

The dried Bark of the stem and branches of cultivated plants of *Cinchona succirubra*, Pav.

The dried Bark of *Cinchona Succirubra* only is official in the *B.P.* and *P.G.* It is official also in the *U.S.P.* under the heading *Cinchona Rubra*, under *Cinchona* is given the dried bark of *Cinchona Ledgeriana*, Howard, *Cinchona Calisaya*, Wedd., *Cinchona officinalis*, L., and of hybrids of these with other species of *Cinchona*. The *B.P.* Bark when used for preparing the official galenical preparations is required to yield from 5 to 6 p.c. of total alkaloids, not less than half of which should consist of Quinine and Cinchonidine, as assayed by the process outlined below. This is not considered an excessive demand, as almost the whole of these mixed alkaloids might consist of Quinine, and it affords no criterion of the amount of Quinine present. It has been pointed out (*P.J.* (3) xvi 407) that a bark may contain the requisite total alkaloids and the official percentage of Quinine and Cinchonidine, and still contain only a trace of Quinine, what, therefore, is really wanted in the *Pharmacopœia* is a Quinine standard for the Bark. The *French Codex* (1908) requires the Bark to contain at least 5 p.c. of total alkaloids and to yield at least 1.5 grammes of crystallised basic Quinine Sulphate containing 8 molecules of Water of crystallisation, this quantity corresponding to 1.257 p.c. of basic Quinine Sulphate, dried at 100° C. (212° F.), or to 1.092 p.c. of anhydrous Quinine. This is the first instance in which a Quinine standard has been adopted by an important Pharmacopœia. An outline of the process adopted for the determination of the alkaloids is too lengthy for inclusion here, but the essential details are given at the end of the *Materia Medica*. The galenical preparations of the *B.P.* are standardised to contain a definite percentage w/v of total alkaloids. The *U.S.P.* *Succirubra* Bark or its hybrids is required to yield not less than 5 p.c. of anhydrous *Cinchona* alkaloids when assayed by the process indicated for the other varieties of the Bark. The other varieties of *Cinchona* bark official in the *U.S.P.* are required to yield not less than 5 p.c. of total anhydrous *Cinchona* alkaloids, of which at least four-fifths shall consist of anhydrous Ether-soluble alkaloids. The galenical preparations of the *U.S.P.* are standardised to contain a definite percentage of Ether-soluble alkaloids.

The *PG* gives no definite requirement, but if the result of the volumetric determination be expressed in terms of the mean combining weights of Quinine and Cinchonidine it should yield not less than 5.07 p.c. of alkaloids. The new *Swiss Ph* adopts a minimum of 6.5 p.c. of alkaloids.

Medicinal Properties—Tonic, bitter, stomachic and astringent. It is valuable in neuralgia and in convalescence from acute diseases, in diarrhoea, excessive perspiration, chronic discharges from mucous membranes, and in dipsomania. (See also *Quinine*.)

Official Preparations—Extractum Cinchonæ Liquidum, Infusum Cinchonæ Acidum, Tinctura Cinchonæ, Tinctura Cinchonæ Composita, and is a source of the Alkaloid Quinine.

Not Official—Decoctum Cinchonæ, Elix. Cinchonæ, Mistura Cinchonæ, Mistura Cinchonæ Acida, Sirop de Quinquina, Tinctura Chinæ Composita, Vinum Chinæ, also Ferratum, Cinchonidinæ Hydrobromidum, Cinchonidinæ Sulphas, Cinchoninæ Iodo Sulphas, Cinchoninæ Sulphas, and Acidum Chinicum.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Descriptive Notes—Red Cinchona bark is now chiefly imported from Ceylon and Java, although some is occasionally offered from the African Island of St. Thomé. The tree grows rapidly, and consequently the bark shinks a good deal in drying, and presents, especially in the branch bark, a shrivelled or longitudinally wrinkled character, it has a reddish colour when broken, and a bitter and astringent taste. Its most marked feature is the presence of reddish warty lenticels. In the variety with leaves hairy beneath, formerly cultivated in Jamaica, these occur very sparingly. These two characters distinguish the red Cinchona from the other barks in use, since the warts are absent in that of *C. Calisaya*, Wedd., and in it the ridges or wrinkles are replaced by longitudinal fissures, which are at first shallow, but become deeper in older barks, and the epidermal layer often exfoliates, showing transverse cracks in the layer beneath. The variety *Ledgeriana*, Howard, is remarkable for its relatively greater thickness as compared with that of other species, but externally resembles *Calisaya*, except that the surface is usually rougher. In *Cinchona officinalis*, Hook., there are numerous transverse cracks with thickened edges, broken into points, so that the bark is rougher to the touch than other kinds. *Cinchona lancifolia*, Mutis, and its varieties are characterised by a whitish spongy epidermal layer and by a loose fibrous fracture. A hybrid of *C. officinalis*, L., and *C. succubra*, Pav. (*Cinchona robusta*, Hort.), presenting the thick-edged cracks of the one and the warty character of the other, is remarkably rich in alkaloid, and is sometimes offered as 'Ledgeriana' bark. Small chips of red barks obtained by spoke-shaving do not present marked characters, and should only be purchased on analysis. The official bark is in quills or somewhat curved pieces coated with periderm, it may vary considerably in length, 2 to 12 in (5 to 30 cm.) or more, and the thickness of the bark itself may vary from about $\frac{1}{10}$ to $\frac{1}{4}$ in ($2\frac{1}{2}$ to 6 mm.), the fracture is shortly fibrous in the smaller, and finely fibrous in the larger pieces, the powder should be brownish or reddish-brown.

Tests Cinchona is one of the few instances in which the *BP* adopts a standard for the drug and indicates a method for the determination of its alkaloidal strength. The process official in the *BP* 1898 is an adaptation of that of the *BP* 1885, which is in turn based upon a process recommended by Squibb, it depends upon the liberation of the alkaloids from the combinations in which they exist in the bark by means of Calcium Hydroxide and their extraction by means of a mixture of 3 parts by volume of Benzol and 1 part by volume of Amyl Alcohol, the alkaloids in turn being shaken out from this solvent by a mixture of Diluted Hydrochloric Acid, and Water, which solution, after careful neutralisation with Ammonia Solution, is concentrated and the Quinine and Cinchonidine precipitated by Sodium Potassium Tartrate Solution, the Tartrates of the remaining alkaloids being precipitated from the filtrate from the Quinine and Cinchonidine by the addition of a slight excess of Ammonia Solution. The *BP* process may be briefly outlined as follows. A weighed quantity (20 grammes) of the finely powdered Bark is treated with 6 grammes of Calcium Hydroxide. The mixture after being moistened with 20 c.c. of Water is well incorporated, and allowed to stand for an interval of one or two hours. It is then transferred to a flask, mixed with 130 c.c. of a mixture of 3 parts by volume of Benzol and 1 part by volume of Amyl Alcohol and extracted by boiling under a reflux condenser for about 30 minutes. The Benzolated Amyl Alcohol is removed, passed through a filter, and the residue in the flask again boiled with a further quantity of the same mixture of Benzol and Amyl Alcohol, the liquid removed as before, and the process repeated a third time if necessary, the residue in the flask being finally transferred to the filter and washed by percolation with the mixture of Benzol and Amyl Alcohol until exhausted of alkaloids. This may be determined by evaporating a few drops of the Benzol-Amyl Alcohol Solution on a watch-glass, acidifying the residue with a drop or two of Diluted Sulphuric Acid, and adding a drop or two of Potassium-mercuric Iodide (Mayer's) Solution. The filtrates are mixed, transferred to a separator, and well shaken whilst warm with a mixture of 2 c.c. of Diluted Hydrochloric Acid and 12 c.c. of Water the liquids are allowed to separate, the acid aqueous solution of the alkaloidal Hydrochlorides is removed and the extraction of the alkaloids remaining in the Benzolated Amyl Alcohol solution accomplished by repeatedly shaking with Water made slightly acid with Hydrochloric Acid. The mixed acid liquids should, whilst warm, be exactly neutralised with Ammonia Solution concentrated to a volume of 16 c.c., and the Quinine and Cinchonidine precipitated as tartrates by the addition of a solution of about 1.5 grammes of Sodium Potassium Tartrate dissolved in 3 grammes of Water, the whole well stirred with a glass rod, and allowed to remain at rest for about an hour. The precipitated Quinine and Cinchonidine Tartrates are filtered, washed, dried at a temperature of the water-oven and weighed. They should contain four-fifths of their weight of alkaloids, and if this quantity be multiplied by 5 the product will be percentage by weight of Quinine and Cinchonidine.

The filtrate from the Quinine and Cinchonidine Tartrates is made slightly alkaline by the addition of Ammonia Solution, the precipitate collected on a filter, washed, dried, presumably also at a temperature of the water-bath, and weighed. This precipitate consists of the alkaloids other than Quinine and Cinchonidine, and if their weight be multiplied by 5 the product represents the percentage by weight present in the Bark, and when added to that of the Quinine and Cinchonidine obtained by the previous process yields the percentage by weight of total alkaloids.

The *BP* process is lengthy and tedious, it requires careful manipulation, and the closest attention to details is necessary to obtain accurate results. The exhaustion of Bark by the hot Benzolated Amyl Alcohol requires considerable time and patience. The exact neutralisation of the Hydrochloric Acid solution of the alkaloids requires great care, as does also the evaporation of the solution to the prescribed volume, and after evaporation the reaction of the liquid should again be ascertained and if necessary again exactly neutralised, unless exactly neutral there will be a liability to loss of alkaloid. It by no means follows that although the liquid has been neutralised before evaporation it will remain neutral during evaporation, and a further addition of Ammonia Solution is generally necessary.

The *USP* employs Ether-chloroform as a solvent for the alkaloids, and divides the process into two parts: (1) for anhydrous Cinchona alkaloids, and (2) for Ether-soluble alkaloids. The outlines of the process are essentially as follows:—A weighed quantity of 15 grammes of Cinchona Bark in No. 80 powder (or finer) is shaken vigorously in an Erlenmeyer flask with a mixture of 250 c.c. of Ether and 50 c.c. of Chloroform, and allowed to stand for 10 minutes. A measured quantity of 10 c.c. of Ammonia Solution is added, and, with frequent intervals of shaking, the mixture is allowed to stand for 5 hours. A measured quantity of 15 c.c. of Water is added, the mixture vigorously shaken and allowed to stand a few minutes. A measured quantity of 200 c.c. of the clear supernatant liquid is then transferred to a separator and the alkaloids extracted by vigorously shaking with 15 c.c. (or sufficient to make the liquid distinctly acid) of Normal Volumetric Sulphuric Acid Solution. The lower acid liquid is drawn off after the two layers have been separated, and the Ether-chloroform solution is again shaken vigorously, with a mixture of 5 c.c. of Normal Sulphuric Acid and 5 c.c. of Water, after separation, the acid liquid is again removed, the shaking is repeated a third time, using 5 c.c. of Water only, and the aqueous liquid is removed. The mixed acid liquids are filtered into a graduated cylinder, the containing vessel and filter are washed with sufficient Water to bring the volume of the contents of the cylinder to 50 c.c., and this measured quantity is then divided into two equal portions of 25 c.c. each. No. 1 quantity of 25 c.c. is placed in a separator, is rendered alkaline by a sufficiency of Ammonia Solution, and the alkaloids removed by shaking carefully for 1 minute with a mixture of 3 volumes of Chloroform and 1 volume of Ether. The lower layer, after separation of the liquids is complete, is drawn off into a tared flask. The

contents of the separator are again shaken with 20 cc of a similar mixture of Chloroform and Ether for 1 minute, and this is followed by a third shaking with 10 cc of the Chloroform-ether mixture, the chloroformic liquids being in each case removed after the liquids have separated into two layers. The mixed Chloroform-ether solutions are evaporated to dryness on a water-bath, the dried residue is mixed with 3 cc of Ether, and again evaporated to dryness. It is then dried in an air-bath at a temperature of 110° C (230° F) until the weight remains constant. This weight multiplied by 20 indicates the percentage by weight of total Cinchona alkaloids. The second quantity of 25 cc is rendered alkaline with a sufficiency of Ammonia Solution, and shaken moderately for 2 minutes with 25 cc of Ether, the temperature of the liquid being maintained below 20° C (68° F), and the liquids allowed to stand for 10 minutes at 15° C (59° F). After separation of the two liquids the aqueous layer is removed and the ethereal solution is transferred to a tared flask. The separator is rinsed out with 5 cc of Ether and the washings added to the main quantity. The Ether is carefully evaporated on a water-bath, the flask and contents dried for 2 hours at a temperature of 110° C (230° F), cooled and weighed. The weight multiplied by 20 yields the percentage by weight of anhydrous Ether-soluble alkaloids. The *USP* adds a note to the effect that the Ether-soluble alkaloids include Quinine, Quinidine, and Cinchonidine.

The above process works well, is easily manipulated and yields the alkaloids in a very fair state of purity. Determinations carried out in the author's laboratory have shown an average of about 5.84 p.c. of total alkaloids and 4.32 p.c. of Ether-soluble alkaloids.

The *PG* describes a method for the determination of the percentage of total alkaloids, but gives no process by which the amount of Quinine and Cinchonidine may be judged. A qualitative test is introduced, which requires that 5 cc of the reserve portion remaining after the quantity for the volumetric determination has been removed, when mixed with 1 cc of Chlorine Water, shall, on the addition of Ammonia Water, yield a fine green coloration.

The process for the quantitative volumetric determination of the alkaloids is as follows:—A weighed quantity of 12 grammes of the finely powdered Bark dried at 100° C (212° F) is mixed in a well-stoppered flask or bottle with 90 grammes of Ether and 30 grammes of Chloroform, a measured quantity of 10 cc of Sodium Hydroxide solution (15 p.c.) is added, and with frequent intervals of vigorous shaking the mixture is allowed to stand for 3 hours. A measured quantity of 10 cc of Water, or sufficient to cause the powdered Bark to sink on shaking, is added, and after the Chloroform-ether solution has separated as a clear layer, a weighed quantity of 100 grammes is filtered through a dry well-covered filter into a flask, and half of the liquid distilled, the remainder is transferred to a separator, the flask is washed out with three successive quantities of 5 cc of a mixture of 3 parts by weight of Ether and 1 part by weight of Chloroform, and the alkaloids are extracted from the mixed Chloroform-ether liquids by agitation with 25 cc of Decinormal Volumetric

Hydrochloric Acid Solution The acid layer is drawn off after the liquids have completely separated and after sufficient Ether has been added to the mixture to cause the Chloroform ether layer to float on the acid liquid, it is filtered through a small paper moistened with Water, into a flask of 100 c c capacity. The extraction of the Chloroform-ether solution is thrice repeated, using 10 c c of Water for each extraction, and the separated aqueous liquids are passed through the same filter paper, the latter is washed with Water, and the mixed filtrates and washings are diluted to 100 c c. A measured quantity of 50 c c is removed, a freshly prepared solution of a small crystal of Hæmatoxylin in 1 c c of Alcohol (90 p c) added, and sufficient Deci-normal Volumetric Potassium Hydroxide Solution added to change the yellow colour to a bluish-violet, the mixture being shaken after each addition. Not more than 4.3 c c should be necessary. No factor is recorded by which the result of the above volumetric test may be calculated into its equivalent in alkaloids. Assuming that the mixture contains equal proportions of Quinine and Cinchonidine, a factor of 0.030931 may be employed, which indicates 5.07 p c of alkaloids. That the alkaloids are not always present in these proportions is evidenced by the observed discrepancies between the results of gravimetric and volumetric determinations. The new *Swiss Ph* states that 1 c c of Deci-normal Volumetric Hydrochloric Acid is equivalent to 30.4 mg of alkaloids.

The use of Hydrochloric Acid for the titration of Quinine instead of Sulphuric Acid eliminates the troublesome fluorescence which is produced when the alkaloid is taken into solution in the latter acid, and which may often seriously interfere with the end reaction. It must be borne in mind that the behaviour of Quinine towards certain indicators of neutrality is somewhat anomalous. With Hæmatoxylin or Cochineal the point of neutrality is reached with the formation of the Hydrochloride ($C_{20}H_{24}N_2O_2 \cdot HCl$), but this salt is alkaline in reaction towards Methyl Orange, and the point of neutrality with this indicator is only reached with the formation of the Acid Hydrochloride ($C_{20}H_{24}N_2O_2 \cdot 2HCl$). Thus, using Hæmatoxylin Solution as an indicator, 1 c c of Deci-normal Volumetric Hydrochloric Acid Solution is equivalent to 0.03218 gramme of anhydrous Quinine, whilst using Methyl Orange Solution 1 c c of the Deci-normal Volumetric Solution is equivalent to 0.01609 gramme of anhydrous Quinine. The behaviour of Cinchonine and Cinchonidine towards these indicators is still more anomalous. It may therefore be doubted whether the application of a purely volumetric method of determination is advisable, and the volumetric result should always be controlled by a gravimetric determination.

The ash varies from 2 to 4 p c.

. Preparations

EXTRACTUM CINCHONÆ LIQUIDUM. LIQUID EXTRACT OF CINCHONA

A dark reddish-brown liquid prepared from Red Cinchona Bark by treatment with Distilled Water acidulated with Hydrochloric Acid

and containing a small proportion of Glycerin, it is officially required to contain 5 p c w/v or 5 grains of Red Cinchona alkaloids in 110 minims, the *USP* Fluid Extract is required to contain 4 p c w/v or 4 grains of anhydrous Ether-soluble alkaloids in 110 minims. A fluid extract does not appear in the *PG*, which, however, has two solid extracts, an aqueous and an alcoholic

Dose—5 to 15 minims = 0.3 to 0.9 cc

22 minims contain 1 grain of alkaloids

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Jap, Mex, Norw, Swed, Swiss and U S, 1 in 1, Solid Extracts—Austri and Hung, Aqueous, Belg, Dutch, Ital, Jap, Mex, Russ, Span and Swiss, Alcoholic, Fr, Ger, Mex and Port, both Aqueous and Alcoholic. Belg has also Chinese Fluidextractum cum Kalio Iodati. Not in the others.

Tests—The specific gravities of commercial fluid extracts of Red Cinchona Bark vary between 1.100 and 1.150, the percentage w/v of total solids from 33.4 to 53.0 p c and the percentage w/v of total alkaloids from 4.6 to 5.46 p c. One commercial fluid extract, purchased as *BP*, assayed in the author's laboratory, possessed a specific gravity of 1.080, contained 27.5 p c w/v of total solids and only 2.6 p c of total alkaloids.

The *BP* adopts a standard of total alkaloids, the *USP* of anhydrous Ether-soluble alkaloids, the former Pharmacopœia requires a mixture of 3 parts by volume of Benzol and 1 part by volume of Amyl Alcohol for the initial extraction of the alkaloids, the latter a mixture of Ether and Chloroform.

The outlines of the official process are essentially as follows.—A measured quantity of 5 cc of the Liquid Extract is diluted with five times its volume of Water, introduced into a separator, rendered alkaline with 15 cc of Potassium Hydroxide Solution, and the liberated alkaloids extracted by well shaking the mixture with 30 cc of a mixture of 3 parts by volume of Benzol and 1 part by volume of Amyl Alcohol, the Benzol-Amyl Alcohol solution is transferred to another separator and the agitation repeated with another 30 cc of a similar mixture, the Benzol-Amyl Alcohol layer being again drawn off into the second separator, the mixed Benzol-Amyl Alcohol liquids are washed with a little Water, rejected, and the alkaloids extracted with 30 cc of a mixture of 1 volume of Diluted Hydrochloric Acid and 5 volumes of Water, the acid solution of the alkaloidal Hydrochlorides is separated, and the extraction repeated with a further 30 cc of a similar mixture of acid and Water. The mixed acid liquids are rendered strongly alkaline with Ammonia Solution, and the liberated alkaloids extracted by agitation with three successive quantities of 10 cc of Chloroform, the chloroformic layers are each in turn separated, mixed, transferred to a weighed dish, the Chloroform is evaporated off, and the residue dried at 110° C (230° F). This weight multiplied by 10 gives the percentage by volume of total alkaloids present in the sample. The *BP* process is stated (*YBP* '05, 362) to give rise to obstinate emulsions, which may be overcome by using 10 cc alcoholic, in place of 15 cc of aqueous, Potassium Hydroxide solution. A suggestion

is also made to weigh instead of to measure the quantity of Liquid Extract

The *USP* process is carried out on the following lines — A measured quantity of 10 c c of the Fluid Extract is introduced into an Erlenmayer flask, rendered alkaline by the addition of 10 c c of Ammonia Solution, and the alkaloids extracted by shaking vigorously for 10 minutes with a mixture of 100 c c of Ether and 25 c c of Chloroform. 66 c c of the clear supernatant liquid is transferred to a separator, the vessel in which the liquid is measured being washed out with 5 c c of Ether, which is in turn added to the contents of the separator. The alkaloids are then removed from the Ether-Chloroform solution by shaking it vigorously for several minutes with a sufficiency (10 c c) of Normal Volumetric Sulphuric Acid Solution, and the acid layer is transferred into another separator. The complete extraction of the alkaloids from the Chloroform-Ether solution is ensured by a further extraction with 5 c c of Normal Volumetric Sulphuric Solution and 5 c c of Water, which is in turn followed by an extraction with 5 c c of Water. The acid aqueous and the aqueous liquids are separated, mixed with the acid layer already contained in the second separator, the temperature of the mixed liquids being maintained below 25° C (77° F), 25 c c of Ether added and sufficient Ammonia Solution to yield an alkaline reaction to red Litmus paper, after vigorous agitation for 2 minutes the temperature is reduced to below 15° C (59° F), and the liquids allowed to stand for 10 minutes at that temperature. The ethereal layer is separated, transferred to a tared flask, the separator washed with 5 c c of Ether, the washings being added to the main quantity, and the Ether evaporated at a moderate heat on a water-bath, the residue being dried for half an hour in an air bath at 110° C (230° F), cooled and weighed, the heating being repeated and the weight taken when constant. The weight of this residue multiplied by 20 gives the percentage w/v of anhydrous Ether-soluble alkaloids present in the Fluid Extract.

The process has been tried in the author's laboratory, and works well, the resulting alkaloids are fairly free from colour, the average percentage of anhydrous Ether-soluble alkaloids in commercial Fluid Extracts was found to be 4.5 p c.

INFUSUM CINCHONÆ ACIDUM ACID INFUSION OF CINCHONA.

Red Cinchona Bark, 1, Aromatic Sulphuric Acid, $\frac{1}{4}$, Distilled Water, boiling, 20, infuse for one hour, and strain (1 in 20)

Dose — $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 c c

Foreign Pharmacopœias — Official in Russ (C Rubra), 1 in 8, with Phosphoric Acid, Fr (Tisanne), 1 in 50, Span, 1 in 33 $\frac{1}{2}$, with Acidum Sulphuricum Alcoholisatum. Not in the others.

TINCTURA CINCHONÆ TINCTURE OF CINCHONA

A clear reddish or brownish-red liquid, prepared by treating red Cinchona bark, in No 40 powder, with sufficient Alcohol (70 p c) to form a tincture which shall contain not less than 0.95 nor more than

1.05 pc w/v of total alkaloids The *USP* Tincture is required to contain 0.75 pc w/v anhydrous Ether-soluble alkaloids The *PG* tincture is calculated to contain about 1 pc w/v of total alkaloids

Dose— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

22 minims contain $\frac{1}{2}$ grain of alkaloids

Foreign Pharmacopœias—Official in Fr, Teinture de Quinquina Rouge, Dan, Noiw, Russ and Swed, Tinctura Chinæ (from any species), Dutch and Ger, Tinctura Chinæ, and Hung, Tinctura Chinæ Simplex (from *C. Succirubra*), Ital, Tinctura di China, Jap, Tinct Chinæ, Mex, Tintura de Quina, Port, Tintura de Quina (from *C. Flava*), Span, Tintura Alcoholica de Quina (from *C. Calisaya* and *C. Loja*), Swiss, Tinctura Cinchonæ, U.S., Tinctura Cinchona (*C. any species not Red*), all 1 in 5, and all by weight, except U.S. Not in Aust. Belg (*C. various species, especially Succirubra*), contains 1 pc w/v total Alkaloids, including at least 0.2 pc Quinine U.S., Cinchona (*USP*), in No. 60, Glycerin, 75, and sufficient of a mixture of Alcohol (95 pc) 675, 250, to produce 1000 by percolation, containing 0.75 pc w/v Ether-soluble Alkaloids

Tests—Tincture of Cinchona has a specific gravity of 0.915 to 0.920, contains from 3.5 to 8.5 pc w/v of total solids and from 62.5 to 66.5 pc w/v of Absolute Alcohol The *BP* requires that 10 cc of Tincture, when assayed to the process described in the large type under *Extractum Cinchonæ Liquidum*, shall yield not less than 0.095 gramme nor more than 0.105 gramme of alkaloids The *USP* standardises the Tincture to a percentage w/v of anhydrous Ether-soluble alkaloids A measured quantity (50 cc) of the Tincture is evaporated to about one-fifth its volume on a water-bath, the liquid transferred to a bottle of about 180 cc capacity, the dish rinsed with 10 cc of diluted Alcohol, and the determination completed as in the case of the fluid extract The weight of anhydrous Ether-soluble alkaloids obtained, multiplied by 4, shows the percentage w/v present in the Tincture

TINCTURA CINCHONÆ COMPOSITA. COMPOUND TINCTURE CINCHONA

A brownish-red liquid possessing an aromatic odour and bitter taste, which is officially required to contain not less than 0.45 nor more than 0.55 pc w/v of the total alkaloids of Red Cinchona Bark, when assayed as described below The Compound Tinctures official in the *USP* and *PG* are prepared from Red Cinchona Bark containing the official percentage of alkaloids required by each Pharmacopœia, but the alkaloidal content is not verified by a determination See also below

Tincture of Cinchona, 20 fl oz, Dried Bitter-Orange Peel, well bruised, 2 oz, Serpentry Rhizome, in No. 40 powder, 1 oz, Cochineal, in powder, 56 grains, Saffron, 110 grains, Alcohol (70 pc), q s to yield 40 fl oz

Made with standardised Tincture of Cinchona instead of the Red Cinchona Bark ordered in the previous edition of *BP*

Dose— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc.

22 minims contain $\frac{1}{2}$ grain of alkaloids.

Foreign Pharmacopœias—Official in Austr, Ger, Hung, Jap and Russ, Tinct Chinæ Comp, also Swiss (Tinct Cinch Co), with Cinchona, Gentian, Orange Peel and Cinnamon (various strengths), Belg (Tinct Whyttii or Tinct Huxham), Cinchona, Orange Peel, Cinnamon and Saffron Dan, Dutch, Norw and Swed (Tinct Chinæ Comp), similar to the above but without Cinnamon, Mex (Tintura de Quina Compuesta), and Span (Tinctura Alcoholica de Corteza de Naranja Compuesta, Tintura corroborante de Whytt), Bitter Orange Peel, Cinchona and Gentian, Port (Tinct de Quina Comp), Cinchona, Orange Peel and Serpentry, U S, Red Cinchona (*USP*), 100, Bitter Orange Peel, 80, Serpentry, 20, Glycerin, 75, and sufficient of a mixture of Alcohol (95 p c) 675, with Water 250, to produce 1000 by percolation Not in Fr

Huxham's Tincture of Bark (Original Formula in 1788)—Powdered Peruvian Bark, 4 oz, Orange Peel, 8 oz, Serpentry Root, 80 grains, Saffron, 160 grains, Cochineal, 80 grains, Brandy, 40 fl oz, digest 3 or 4 days

Tests—Compound Tincture of Cinchona possesses a specific gravity of 0.910 to 0.920, contains about 5 p c w/v of total solids and about 63.0 p c w/v of Absolute Alcohol. A measured quantity of 10 cc, when assayed by the official process as outlined in the large type under *Extractum Cinchonæ Liquidum*, should yield an amount of alkaloids corresponding to not less than 0.45 nor more than 0.55 p c w/v of total alkaloids. Neither the *USP* nor the *PG* gives a process for the determination of the alkaloids, the presence of the soluble principles from the other constituents of the Tincture invalidating the results.

The residue obtained by the evaporation to dryness of 2 cc of the *BP* Compound Tincture should impart a yellow colour to Chloroform, and if the chloroformic solution be filtered and evaporated to dryness, the residue when moistened with a drop of concentrated Sulphuric Acid will acquire a beautiful indigo-blue tint, indicating the presence of Saffron. The *USP* and *PG* Compound Tinctures are made from Red Cinchona Bark, and do not contain Saffron.

Not Official.

DECOCTUM CINCHONÆ—Red Cinchona Bark, in No. 20 powder, 1½, Distilled Water, 20, boil 10 minutes, when cold, strain, and pour on the marc sufficient Water to make 20 (1 in 16)

Dose—1 to 2 fl oz = 28.4 to 56.8 cc

Official in Belg, 1 of Fluid Extract in 10, Dan, 1 in 8 with Hydrochloric Acid, Dutch, 6 in 100, Ital, 1 in 20, Norw and Swed, 1 in 10 with Hydrochloric Acid, Port, Cinchona Flava 1 in 10, also Fusca 1 in 10, Russ, Cinchona Rubra, 1 in 7.5, containing Sulphuric Acid, Span, Cocimento de Quina Calisaya and Cocimento de Quina de Loja, each about 1 in 66, also Cocimento de Quina y Valeriana and Cocimento Antiseptico, Dan and Norw have a Dec Chinæ c Senega

ELIXIR CINCHONÆ (*USNF* 1896)—Tincture of Cinchona, 12, Syrup, 10, Glycerin, 10, Aromatic Elixir, 48. Each fl oz represents about 14 grains of Yellow Cinchona. A similar preparation is made with Detannated Tincture of Cinchona for use in combination with preparations of Iron.

This has been adopted by the *BPC*, but in the latest edition of *USNF* (1906) the preparation is made with the alkaloids, and not with Tincture of Cinchona.

MISTURA CINCHONÆ (*for children*)—Diluted Nitric Acid, 30 minims, Tincture of Cinchona, 2 fl drms, Glycerin, 1 fl drms, Distilled Water, to 1½ fl oz
—*Middlesea*

Dose—1 to 2 fl drms

MISTURA CINCHONÆ ACIDA—Liquid Extract of Cinchona, 10 minims, Diluted Nitric Acid, 10 minims, Aromatic Syrup, 80 minims, Water, to 1 fl oz —*St Thomas's*

This formula has been incorporated in the *B P C*

Liquid Extract of Cinchona, 10 minims, Diluted Nitric Acid, 10 minims, Aromatic Syrup, 60 minims, Water, to 1 fl oz —*Brompton*

Other hospitals include *Mistura Cinchonæ Acida*, but they are made with *Decoctum Cinchonæ*

SIROP DE QUINQUINA—Percolate 1000 of Red Cinchona in No 26 powder with 1000 of Alcohol (80 p c), displacing with Water to obtain 1000 of percolate, distil off 445 and dissolve in the residue when cold 1000 of Sugar —*Fr.*

TINCTURA CHINÆ COMPOSITA—Cinchona Bark, 6, Orange Peel, 2, Gentian Root, 2, Cinnamon Bark, 1, Diluted Spirit (60 p c), 50 —*Ger*

VINUM CHINÆ (*Ger* and *Jap*)—Dissolve Gelatin, 1, in warm Water, 10, mix with Sherry, 1000, add powdered Cinchona Bark, 40, allow to stand for eight days at 15° to 20° C Press, and to the expressed liquor add Sugar, 100, Tincture of Orange, 2, allow to stand in a cool place for fourteen days, and filter.

Austr, Vinum Chinæ—Dissolve 1 of Gelatin in 20 of boiling Water, and mix with 780 of Malaga Wine, after 24 hours, add 50 of Fluid Extract of Cinchona, 50 of Tincture of Orange, and 100 of Clarified Honey

Belg, Chinæ Vinum—1 of Fluid Extract in 50 of stronger Wine

Fr, Vin de Quinquina Officinal—Cinchona, 25, Alcohol (60 p c), 75, Dilute Hydrochloric Acid, 2, Red Wine, 920

Dutch, Vinum Chinæ—1 Cinchona Succirubra percolated with a mixture of 1 of diluted Alcohol, 4 of Malaga Wine, and 8 of Water *g s* to produce 40, in which dissolve 10 of Sugar

Hung, Vinum Chinæ—Extract of Cinchona, 1, Malaga Wine, 80, Simple Tincture of Cinchona, 20

Ital, Vino Chinato—Cinchona, 1, Marsala Wine, 30

Mex, Vino de Quina—Cinchona, 8, Sherry Wine, 100

Norw, Vinum Chinæ—Cinchona, 50, Citric Acid, 1, Alcohol, 20, Malaga Wine 1000

Russ, Vinum Chinæ—Tincture of Cinchona, 1, Sherry Wine, 4

Port, Vino de Quina—Cinchona flavum, 1, Port Wine, 20, Vinho de Quina Cinzenta, Cinchona fuscum, 1, Madeira Wine, 10, Vinho de Quina Composto, Cinchona flavum, 4, Gentian, 1, Bitter-Orange Peel, 1, Port Wine, 100, Vinho de Quina Ferruginoso, Iron and Potassium Tartrate, 1, Vinho de Quina Cinzenta, 200

Span, Vino de Quina—Loja Bark, 1, Sherry Wine, 16 6, Vino de Quina Ferruginoso, Crystallised Ferrous Sulphate and Citric Acid, of each, 1, Distilled Water, 10, Quinine Wine, 500

Swiss, Vinum Cinchonæ—80 of Fluid Extract, 20 of diluted Alcohol, 40 of Milk, 1 of Citric Acid, 910 of Malaga Wine

B P C, Vinum Cinchonæ—Elixir Cinchona, 1, Detannated Sherry, to make 8

Dutch, Vinum Chinæ Feriatum—1 of Ferri *r* h Ammonium Citrate, dissolved in 4 of Water, and added to 95

Austr, Vinum Chinæ Feriatum—1 of Gelatin, dissolved in 20 of boiling Water, is added to 955 of Malaga Wine, after 24 hours, add 5 of Iron and Quinine Citrate dissolved in 20 of Water

Norw Vinum Chinæ Feriatum—Iron and Ammonium Citrate, 1, Cinchona Wine, 100

Swed, Vinum Chinæ Feriatum—Iron and Quinine Citrate, 1, Syrup, 10, Marsala Wine, 89

B P C, Vinum Cinchonæ Ferratum.—Iron and *r* h C, 1, Cinchona Wine, 200

CINCHONIDINÆ HYDROBROMIDUM ($C_{10}H_{12}N_2O$, HBr , H_2O , eq. 990 28) —Long, light yellow, odourless prismatic crystals, possessing a very bitter taste It contains 71 23 p c of Cinchonidine and 4 58 p c of Water Soluble 1

in 40 of Water Under the name of 'Blennostasine,' a combination similar to this has been introduced for the treatment of affections characterised by catarrhal hypersecretion

Tests—An aqueous solution of the salt yields with Potassium Sodium Tartrate solution a white precipitate soluble in diluted Hydrochloric Acid, with Ammonia Solution it yields a white precipitate soluble in Chloroform The aqueous layer when separated from the Chloroform and rendered faintly acid with Nitric Acid yields with Silver Nitrate Solution a yellowish curdy precipitate soluble with difficulty in Ammonia Solution, insoluble in Nitric Acid

When ignited with free access of air 0.5 gramme of the salt should leave no weighable residue

An aqueous solution should yield no precipitate on the addition of a solution containing a soluble Sulphate

CINCHONIDINÆ HYDROBROMIDUM ACIDUM ($C_{16}H_{19}N_2O$, $2HBr$, $2H_2O$, eq. 488.51)—Light yellow prismatic crystals, readily soluble in Water It should contain 59.78 p.c. of Cinchonidine and 7.32 p.c. of Water

Tests—An aqueous solution of the acid salt responds to tests described under Cinchonidine Hydrobromide

The aqueous solution should yield no precipitate on the addition of a solution of a soluble Sulphate

When ignited with free access of air 0.5 gramme of the salt should leave no weighable residue

CINCHONIDINÆ SULPHAS ($C_{16}H_{19}N_2O$, H_2SO_4 , $3H_2O$, eq. 735.08)—Colourless, odourless, silky, acicular crystals, having a very bitter taste It contains 79.46 p.c. of Cinchonidine and 7.29 p.c. of Water

In doses of one and a half to twice those of Quinine, is stated to form a reliable substitute and frequently to be better tolerated —*P*, lxxiii 682

Solubility—1 in 100 of Water, 1 in 60 of Alcohol (90 p.c.), insoluble in Chloroform and Ether

Dose—1 to 10 grains = 0.06 to 0.65 gramme

Foreign Pharmacopœias—Official in Fr. and U.S. Not in the others

The salt is capable of forming a number of Hydrates, according to the strength of solutions from which it is crystallised From a moderately concentrated solution it crystallises with $6H_2O$, from a hot concentrated solution it crystallises with $3H_2O$ The formula for the salt official in the *French Codex* (1908) shows 6 molecules of Water of crystallisation, which is equivalent to 13.6 p.c. of Water It contains 74.06 p.c. of Cinchonidine

Tests—Cinchonidine Sulphate loses its Water at $100^\circ C$ ($212^\circ F$), and the anhydrous salt reabsorbs moisture on exposure to moist air It has a melting point of 205° to $206^\circ C$ (401° to $402.8^\circ F$), slightly below which temperature it darkens in colour An aqueous solution of the salt is neutral in reaction towards Litmus paper, and yields on the addition of Ammonia Solution a white precipitate, only slightly soluble in an excess of the reagent, but soluble in Ether, a portion of the salt subsequently crystallising out Another portion of the aqueous solution, acidified with diluted Hydrochloric Acid, yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric or Nitric Acids

The more generally occurring impurities are excess of Water, readily carbonisable organic impurities, Quinine or Quinidine Sulphates, Cinchonine Sulphate, and mineral matter Excess of Water is readily detected by the loss of weight of the specimen at a temperature of $100^\circ C$ ($212^\circ F$) The theoretical percentage of Water in the $3H_2O$ salt is 7.29 p.c. The salt should not lose more than 8 p.c. when dried at this temperature The $6H_2O$ salt of the *Fr. Codex* (1908) loses 13.6 p.c. Readily carbonisable organic impurities may be detected by the behaviour of the salt when treated with concentrated Sulphuric Acid, a pure specimen should not become more than faintly coloured, Quinine or Quinidine Sulphates may be detected by the marked fluorescence produced in a 1 in 1000 solution of the salt by diluted Sulphuric Acid, not more than a faint blue fluorescence should be noticed, Cinchonine Sulphate may be detected by precipitating the Cinchonidine as an insoluble Tartrate and testing the filtrate with

a drop of Ammonia Solution A weighed quantity of 0.5 gramme is macerated at 15° C. (59° F.), with 20 c.c. of Water, 0.5 gramme of Potassium Sodium Tartrate is added and the maceration continued with intervals of 10 minutes for 1 hour, the temperature being maintained at 15° C. (59° F.) A drop of Ammonia Solution added to the filtered liquid should not produce more than a slight turbidity. This test also serves to detect Quinidine Sulphate. Mineral matter may be detected by the residue left on ignition, the salt should leave no weighable residue when ignited with free access of air.

CINCHONINÆ IODO-SULPHAS (Antiseptol) —A brown, or reddish-brown, odourless powder, insoluble in Water, soluble in Alcohol (90 p.c.), and in Chloroform. It contains about 50 p.c. of Iodine. Introduced as a substitute for Iodoform. Used in the form of a 1 in 8 ointment for lupus. Has also been given internally in doses of 1 to 5 grains = 0.06 to 0.32 gramme.

CINCHONINÆ SULPHAS ($C_{19}H_{22}N_2O_2$), $H_2SO_4 \cdot 2H_2O$, eq. 717.20 —Hard, white shining odourless, prismatic crystals, having a very bitter taste.

It is the sulphate of an alkaloid obtained from various species of Cinchona Bark. B.P. 1885 said from Cinchona and Remijna Bark. It contains theoretically 81.44 p.c. of Cinchonine and 4.98 p.c. of Water.

Solubility —1 in 70 of Water, 1 in 9 of Alcohol (90 p.c.), 1 in 60 of Chloroform, sparingly in Ether.

Dose —1 to 10 grains = 0.06 to 0.65 gramme.

Foreign Pharmacopœias —Official in Mex., Port. and U.S. Not in the others.

Tests —Cinchonine Sulphate when heated to 100° C. (212° F.) readily loses its Water of Crystallization, when rendered anhydrous at this temperature it melts at about 150° C. (302° F.). Its aqueous solution is neutral in reaction towards Litmus paper, and is dextrorotatory, and yields on the addition of Ammonia Solution a white precipitate. The aqueous solution when acidified with diluted Sulphuric Acid gives a white precipitate. Water added yields no green coloration on the addition of an excess of Ammonia Solution. The addition of Barium Chloride solution to an aqueous solution of the salt yields a white precipitate insoluble in Hydrochloric Acid. One part of the powdered anhydrous Sulphate should dissolve in 80 parts by weight of Chloroform.

The more generally occurring impurities are excess of moisture, Quinine or Quinidine or Cinchonidine Sulphate, readily carbonisable organic impurities and mineral matter. Excess of moisture is readily shown by the loss in weight of 1 gramme of the salt at 100° C. (212° F.) which should not amount to more than 5.0 p.c., the theoretical percentage as above indicated being 4.98 p.c., Quinine and Quinidine Sulphates produce a marked blue fluorescence in a 1 in 1000 solution of the salt in diluted Sulphuric Acid. Cinchonidine Sulphate is detected, and additional confirmation of the presence of Quinine is detected when 1 part of the anhydrous salt is treated with 80 parts by weight of Chloroform, anhydrous Cinchonine Sulphate will dissolve, anhydrous Cinchonidine Sulphates are insoluble in Chloroform, readily organic impurities may be detected by concentrated Sulphuric Acid, should not yield more than a pale yellow coloration, mineral matter may be detected by the residue left on ignition, the pure salt leaving no residue on incineration.

Cinchonidinæ Sulphas Acidus and Cinchoninæ Sulphas Acidus are also known. These salts are more readily soluble in Water.

Cinchonidinæ Salicylas and **Styracis Salicylas** have been used, the former as a tonic and antiperiodic, the latter as a prophylactic against malaria.

ACIDUM CHINICUM Chinic Acid, Kinic Acid, Quinic Acid, $C_7H_8O_6$, eq. 190.65 —Colourless, transparent, rhombic prisms or flat, crystalline masses, having a strongly acid but not a bitter taste.

Solubility —1 in $2\frac{1}{2}$ of Water, 1 in 42 of Alcohol (90 p.c.), insoluble in Ether.

Tests —Quinic Acid possesses a melting point of 161° to 162° C. (321.8° to 323.6° F.). Its aqueous solution is laevogyrate. A little of the powder

distilled with Manganese Dioxide and Sulphuric Acid yields Quinone, which condenses on the cool side of the tube in the form of deep yellow prisms. The acid may be titrated with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator, a weighed quantity of 1 gramme of the acid should require about 5.2 c.c. of the Volumetric Solution corresponding to about 99 p.c. of pure Chinic Acid. It should leave no residue when ignited with free access of air.

It has been introduced in the treatment of the Uric Acid diathesis and in gout. It is more generally employed in the form of a salt, e.g., Lithium Quinate, *q.v.p.* 738.

Dose — 1 to 5 grains = 0.06 to 0.32 gramme

CINNAMOMI CORTEX.

CINNAMON BARK

FR, CANNELLE DE CEYLAN, GER, CHINESISCHER ZIMMT (CASSIA), ITAL, CANNELLA DEL CEYLAN, SPAN, CANELA DE CEYLAN

The dried inner Bark of shoots from the truncated stocks of *Cinnamomum Zeylanicum*. The bark from cultivated trees is alone official. Imported from Ceylon, and commercially known as Ceylon Cinnamon.

Medicinal Properties — Calminative, astringent, aromatic stimulant, and antiseptic, chiefly used as an adjuvant to other medicines, and as a flavouring agent. Often employed with Chalk in diarrhoea. An essence has been used as a prophylactic against influenza.

60 grain doses for dysentery — *B.M.J.* '95, 1 530, *L.* '95, 1 567. Has been lauded for cancer, but the majority of evidence is not in its favour — *M.A.* '95, 163.

Inhalation of Oil of Cinnamon in the treatment of consumption — *B.M.J.* '96, 11 1374.

The vapour of the Oil of Cinnamon exerted no retarding or inhibitive influence on the growth of the tubercle bacillus — *B.M.J.* '99, 1 203.

Dose — 10 to 20 grains = 0.65 to 1.3 gramme in powder

Official Preparations — Of the Bark, Aqua Cinnamomi, Oleum Cinnamomi, Pulvis Cinnamomi Compositus, and Tinctura Cinnamomi, used in the preparation of Decoctum Hæmatoxyli, Pulvis Catechu Compositus, Pulvis Cretæ Aromaticus, Pulvis Kino Compositus, Tinctura Cardamomi Composita, Tinctura Catechu, and Tinctura Lavandulæ Composita. Of the Water, Mistura Cretæ, Mistura Guaiaci, Mistura Olei Ricini, Mistura Spiritus Vini Gallici, Syrupus Aromaticus and Syrupus Cascariæ Aromaticus. Of the Oil, Spiritus Cinnamomi. Of the Compound Powder, Pilula Aloes et Ferri and Pilula Cambogiæ Composita. Of the Spirit, Acidum Sulphuricum Aromaticum.

Not Official — Pulvis Aromaticus, Tinctura Cinnamomi Composita, Tinctura Aromatica.

Foreign Pharmacopœias — Official in Austri, Dan, Dutch, Fr (Cannelle), Ital (Cannella), Mex (Canela), Norw, Port (Canella), and Swed use Ceylon Cinnamon only. Ger, Hung, Jap and Russ use Chinese Cinnamon or Cassia only. Belg, Span, Swiss and U.S. use both kinds.

Descriptive Notes — By the name of Cinnamon in this country the bark of *Cinnamomum Zeylanicum*, Breyn, imported from Ceylon, is understood. In Germany the Cortex Cinnamomi, official in the P.G., is the bark known as Cassia in this country, but as Chinese

Cinnamon in Germany and in the United States. In the *USP* the Ceylon Cinnamon is official as *Cinnamomum Zeylanicum*, but instead of Cassia or Chinese Cinnamon there is also official a kind known as Saigon Cinnamon, which is the bark of an undetermined species of *Cinnamomum*. Of the Cinnamon tree about six varieties are cultivated in Ceylon, and the different grades are distinguished by being packed in rolls of quills of different sizes, the more slender the rolls the better the quality, therefore the limit of size given in the *BP* $\frac{3}{8}$ inch (9 mm) in diameter indicates the quality intended. The scrapings of the tips of the shoots and the broken fragments of quills have been regularly imported into Europe since 1867 under the name of 'Cinnamon chips' and have been used for the distillation of the oil. The thick trunk bark is sometimes offered in commerce, but has very little aroma, and probably finds its way into the cheaper kinds of mixed spice. Cassia bark in powder is sometimes substituted for that of Cinnamon, but it may be detected by the different flavour, and under the microscope by the presence of cork cells, which are absent from Cinnamon, by the larger and broader bast fibres, and the larger starch grains. Cassia bark is known to drug brokers under the name of 'Cassia lignea' to distinguish it from 'Cassia vera,' which is a hard mucilaginous bark derived from *Cinnamomum Burmanni*, DC, with an allied but different odour, and apparently imported from Padang in Sumatra. Chinese Cassia occurs in small quills 2 to 3 inches (5 to 7.5 cm) long made into packets of about 12 inches (30 cm) long and 3 inches (7.5 cm) in diameter, Cassia vera in quills 12 to 15 inches (30 to 37.5 cm) or more in length, Saigon Cassia, which goes chiefly to the United States, occurs in quills 3 to 4 inches (7.5 to 10 cm) long, and has a more intensely sweet taste and stronger flavour. The *PG* directs that Cassia should not have a mucilaginous taste and that the medullary rays are usually only 2 cells thick, thus excluding the Padang Cassia vera. Other species of Cassia imported into this country may be distinguished under the microscope even in powdered form, see *Museum Report, Pharm Soc* 1903, pp. 50, 51. Some commercial samples of powdered Cinnamon apparently contain a large proportion of Cinnamon bark that has been distilled.

Tests.—Cinnamon Bark leaves on incineration about 4 p.c. of ash, and 6 p.c. is seldom exceeded. Eight samples examined in the author's laboratory gave from 2.8 to 4.26 p.c., with an average of 3.5 p.c., 6 samples of the powder gave from 4.32 to 5.2 p.c., with an average of 4.74 p.c. The *USP* gives not over 4 p.c. No limit of ash is given in the *BP*.

Preparations

AQUA CINNAMOMI. CINNAMON WATER

CINNAMON BARK, bruised, 1, Water, 20, distil 10 (1 in 10)

The distilled 'Aqua' is very turbid from suspended Oil. There is no recognised rule in dispensing as to whether it should be filtered or not, but it is customary to do so.

Dose —1 to 2 fl oz = 28 4 to 56 8 c c

Foreign Pharmacopœias —Official in Austr, Belg, Dan, Dutch, Ger, Jap, Russ, Swed and Swiss, 1 in 10, Fr (Eau de Cannelle), 1 in 5, Ital (Acqua dist di Cannelle) Mex (Agua destilada de Canela), 1 in 4, Hung, 1 in 5, also Aqua Cinnamomi Spirituosa, 3 in 10, Port, 1 in 8, Span (Agua destilada de Canela) C 270, Water 2000, Alcohol (90 p c) 100, distil 1200 Norw and U S, made with Oil 1 in 500

Tests —An approximate idea of the amount of oxidation of the Cinnamic Aldehyde which has occurred may be obtained by acidifying a measured quantity with 25 p c Sulphuric Acid Solution and adding Deci-normal Volumetric Potassium Permanganate Solution till the fluid acquires a pink coloration, remaining permanent for several seconds. The amount of Cinnamic Acid may be determined by titration with Deci-normal Volumetric Sodium Hydroxide Solution. 10 c c of a freshly Distilled Water will require from 12 to 14 c c of the Deci-normal Volumetric Potassium Permanganate Solution, and about 0 1 c c of the Deci-normal Volumetric Sodium Hydroxide Solution, a Water which has been distilled and stocked for some time may require only 4 c c of the Permanganate Solution and about 1 5 c c of the Deci-normal Volumetric Sodium Hydroxide Solution.

OLEUM CINNAMOMI OIL OF CINNAMON

FR, ESSENCE DE CANNELLE DE CEYLAN, GER, ZIMMTEL (CASSIA), ITAL, ESSENZA DI CANNELLA, SPAN, ESENCIA DE CANELA

A light yellow liquid, obtained by distillation from Cinnamon Bark, and possessing the agreeable, delicate, aromatic odour of the Ceylon Cinnamon, and a spicy, sweet, burning taste. It darkens in colour by exposure to light and air. It should be kept in dark amber-tinted, well-closed glass bottles, and protected as far as possible from the air and light.

It usually contains 65 to 75 p c of Cinnamic Aldehyde, from 4 to 8 p c of Eugenol, and some Phellandrene. Yield of oil is about 0 5 to 1 p c.

Cinnamic Acid, an oxidised product of the oil, is described under Acidum Cinnamicum, p 39.

Solubility —10 in 3 of Alcohol (90 p c), 1 in 45 of Alcohol (60 p c)

Medicinal Properties —Possesses the aromatic and antiseptic properties of Cinnamon Bark, without its astringency. It is a powerful local stimulant when administered internally.

Of late years the medicinal virtues of Cinnamon have received a good deal of attention. The Oil has already been used as an inhalation in phthisis. It has been shown in the Bradshaw Lecture on the treatment of enteric fever (*B M J* '04, ii 1451, '05, i 414) that 2½ minims doses given at the commencement, followed by increasing doses up to 5 minims, have given favourable results. An appreciable, though slight, inhibitory influence on the growth of typhoid bacillus begins to be exerted by the oil in a dilution of about 1 in 2600, and when the strength approaches 1 in 1000 its antiseptic effect is complete. The quality of the drug must be above reproach.

Dose —½ to 3 minims = 0 03 to 0 18 c c

In pill or on Sugar

Foreign Pharmacopœias—Official in Dutch, Fr, Ital, Mex (Acete Volatil de Canela), Port and Span use Oil of Cinnamon, Dan, Ger, Hung, Jap, Norw, Swiss and U S use Oil of Cassia Austr and Swed, Cinnamalum (Cinnamic Aldehyde) in place of Oil of Cassia, U S, Cinnaldehydum as well as Oil of Cassia, Belg, Oils of Cinnamon and of Cassia

Tests—Cinnamon Oil should possess a specific gravity of from 1.023 to 1.040, the official figures are from 1.025 to 1.038. Optically it is almost inactive, the rotation not varying more than 1° in a tube of 100 mm. It should dissolve to form a clear solution in 2 parts by weight of Alcohol (70 p c). It is officially required to show the absence of more than 50 p c of non-aldehydic constituents as determined by well shaking a measured quantity of 10 c c with 5 times its volume of a boiling Sodium Hydrogen Sulphite Solution (30 p c), the oily layer which separates being required to measure when cooled to 15.5°C (60°F), not more than 5 c c. The official test is unsatisfactory, and the directions quite inadequate. The Sodium Hydrogen Sulphite Solution should be added in small portions at a time, and care should be taken to heat after each addition until the solid compound liquefies. The oil official in the *BP* is that derived from Ceylon Cinnamon, that of the *USP* and *PG* from Cassia Cinnamon. The *BP* oil should contain not more than 50 p c of non-aldehydic constituents, the *USP* not less than 75 p c, and the *PG* not less than 70 p c of Cinnamic Aldehyde. A determination of the percentage of Cinnamic Aldehyde present in a specimen may be made by the process given under the tests for Oil of Cassia.

The chief adulterant of Cinnamon Oil is oil distilled from Cinnamon leaf. It may be qualitatively detected by dissolving a measured quantity of the oil in 5 times its volume of Alcohol (90 p c), and adding a few drops of Ferric Chloride Test-solution, no decided blue coloration should be produced. If present in quantity it would cause an increase in the specific gravity, the percentage of Cinnamic Aldehyde would be lowered, and the Eugenol content increased. The percentage of Eugenol may be determined approximately by treating the oil with a 5 p c Potassium Hydroxide solution and measuring the diminution in volume. If more accurate results are desired, the Eugenol may be determined by conversion into Benzoyl-eugenol as described in the tests under *Oleum Caryophylli*.

The tests of the *USP* and *PG* are compared under Oil of Cassia.

PULVIS CINNAMOMI COMPOSITUS. COMPOUND POWDER OF CINNAMON *BP Syn*—PULVIS AROMATICUS

Cinnamon Bark, 1, Cardamom Seeds, 1, Ginger, 1, all in powder (1 in 3)

Dose.—10 to 40 grains = 0.65 to 2.6 grammes

Foreign Pharmacopœias—Official in Port (Pó de Canella Comp), Cinnamon 7, Cardamoms 7, Ginger 6, Pulvis Aromaticus, Dutch same as Brit, Swiss, Cinnamon 1, Cardamoms 1, Ginger 1, Sugar 7, U S, Cinnamon 7, Ginger 7, Cardamoms 3, Nutmeg 3. Not in the others.

SPIRITUS CINNAMOMI. SPIRIT OF CINNAMON

Oil of Cinnamon, 1, Alcohol (90 p c), *q s* to yield 10

B P 1885 was 1 in 50

Dose—5 to 20 minims = 0.3 to 1.2 c c

Foreign Pharmacopœias—Official in Belg, 1 in 100, Jap, Cassia Oil 1 in 50, U S, 1 in 10, Dutch, Ital, Mex and Port (distilled from the Bark). Not in the others

Tests—Spirit of Cinnamon has a specific gravity of from 0.850 to 0.855, it leaves about 0.24 p c w/v of residue on evaporation over a water-bath, this residue, when dissolved in Alcohol (90 p c) yields a green coloration on the addition of Ferric Chloride Test-solution

TINCTURA CINNAMOMI TINCTURE OF CINNAMON

1 of Cinnamon Bark, in No 40 powder, percolated with Alcohol (70 p c), to yield 5

B P 1885 was 1 in 8, with Rectified Spirit

Dose— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S, 1 in 5, all by weight except U S

Tests—Tincture of Cinnamon has a specific gravity of from 0.898 to 0.906, contains from 1.8 to 3.0 p c w/v of total solids and from 65 to 69 p c w/v of Absolute Alcohol

Not Official.

PULVIS AROMATICUS (*B P* 1864)—Cinnamon 4, Nutmeg 3, Saffron 8, Cloves $1\frac{1}{2}$, Cardamoms 1, Refined Sugar 25

This has been incorporated in the *B P C* as follows —

Pulvis Aromaticus Compositus—Cinnamon Bark 10.66, Nutmeg 8, Saffron 8, Cloves 4, Cardamom Seeds 2.66, Refined Sugar 66.66

TINCTURA CINNAMOMI COMPOSITA (*P L*)—Cinnamon, 1 oz, Cardamoms, $\frac{1}{2}$ oz, Long Pepper, $2\frac{1}{2}$ drm, Ginger, $2\frac{1}{2}$ drm, Proof Spirit, 40 fl oz

This has been incorporated in the *B P C* as follows —Cinnamon Bark, bruised, 2.50, Cardamom Seeds, bruised, 1.25, Long Pepper, bruised, 1, Ginger, bruised, 1, Alcohol (60 p c), sufficient to produce 100

Port (*Tinctura di Canella Composta*), Cinnamon 10, Cardamoms 4, Cloves 4, Ginger 4, Alcohol (85 p c) 100

TINCTURA AROMATICA (*Ger, Russ and Swiss*)—Cinnamon Root, in coarse powder, 5, Ginger, 2, Galangal Root, 1, Cloves, 1, Cardamoms, 1, Diluted Alcohol, 50

Ger use 60 p c, *Russ* 70 p c, and *Swiss* 68 to 69 p c Alcohol

Austr, same form but with Zedoary Root in place of Galangal Root

Dan and *Norw*, Cinnamon, 4, Ginger, Galangal Root, Cloves and Cardamoms, of each 1, Alcohol (68 p c), 40

Jap, Cloves 2, Cinnamon 10, Cardamoms 2, Ginger 5, Alcohol (68 p c) 100, extract in the cold for 7 days, press, filter, to the filtrate add Spirit of Lemon 5

COCÆ FOLIA.

COCA LEAVES

FR, FEUILLE DE COCA, GER, COCABLATTER, ITAL, FOGLIA DI COCA, SPAN, COCA DEL PERU (HOJADE)

The dried leaves of *Erythroxylum Coca*, and its varieties

Coca leaves contain an amount of alkaloids varying from 0 to 1.5 p c. The average amount is about 0.5 p c. The leaves frequently contain very little alkaloid, owing to the alkaloids readily undergoing decomposition when the leaves are exposed to heat and moisture. The amount of Cocaine in a good sample of leaves is about 70 p c or even less of the total alkaloids. The leaves official in the *U S P* are required to yield not less than 0.5 p c of the Ether-soluble alkaloids of Coca, those official in the new *Swiss Ph* a minimum content of 0.7 p c of alkaloids.

Medicinal Properties—A nerve and muscular tonic, stimulant and restorative. Useful during convalescence, in debility and nervous exhaustion, and to prevent fatigue. The leaves are chewed by the natives of Peru and Bolivia to sustain them during the day, that they may defer eating till the evening.

It has been recommended for the cure of the craving for Opium and for Alcohol, but the craving for Cocaine, which is acquired by the excessive use of Coca, is possibly worse than either.

Official Preparations—*Extractum Cocæ Liquidum*. Used in the preparation of Cocaine and Cocainæ Hydrochloridum.

Not Official.—*Elixir Cocæ*, *Extractum Cocæ*, *Tinctura Cocæ*, and *Vinum Cocæ*.

Foreign Pharmacopœias—Fr, Ital, Jap, Port and U S (*Coca*), Mex and Span (*Coca del Peru*), Swiss (*Folium Cocæ*). Not in the others.

Descriptive Notes.—The Coca leaves of commerce are of three varieties. The Bolivian or Huanuco, the Peruvian or Truxillo, and the Coca leaves cultivated in Java. The first are derived from *Erythroxylum Coca*, Lamarck, and the second from a plant which has been named by Rusby, *Erythroxylum Truxillense*. The third is described by botanists as derived from *E Spruceanum*, Burck. The Bolivian leaves are oval, dark olive green when fresh, with a dark mid-rib, and are usually not broken. The Peruvian are thinner, pale green, oblanceolate and narrower, and are generally much broken. The Java leaves are more lanceolate, darker green, and the mid-rib is reddish towards the base. The leaves as they arrive in commerce vary in size and quality. In the *B P* the size of the Bolivian is defined as $1\frac{1}{2}$ to 3 inches (37 to 75 mm) long and 1 to $1\frac{1}{2}$ inches (25 to 37 mm) in breadth, and oval, but the Peruvian leaves are only described as smaller, narrower, and more brittle than the Bolivian. They are, however, different in shape, being more or less oblanceolate and more tapering towards the base. The Java leaves are not described in the *B.P*. In the Bolivian Coca the mid-rib has a prominent ridge on its upper surface which is not present in the Peruvian. The lateral lines, formed of collenchyma, where

the leaf is folded when young, are more prominent in the Bolivian than in the other varieties. Under the microscope the characteristic features are the papillose polygonal cells of the under surface, the papillæ presenting the appearance of a circle in the centre of each cell, the pericyclic fibres, the prismatic crystals, and the stomata between two narrow cells parallel with the guard cells. Coca leaves for use in pharmacy should not smell or taste mouldy, and should produce a slightly numbing effect on the tongue. They lose alkaloids if exposed to damp, or if not carefully dried.

Tests—Coca Leaves of the *BP* are not required to yield any definite percentage of alkaloids, and no method of determination is given. The *USP* method is carried out on the following lines—A weighed quantity of 10 grammes of the leaves in No 60 powder is transferred to an Erlenmeyer flask and allowed to soak for 10 minutes in 50 c c of a mixture of 1 part by volume of Chloroform and 4 parts by volume of Ether. After the addition of a mixture of 2 c c of Ammonia Solution and 3 c c of Water, the flask is set aside for one hour, with intervals of frequent shaking. The contents of the flask are then transferred to a small percolator having a Cotton-Wool plug packed in the neck and connected with a separator containing a mixture of 6 c c of Normal Volumetric Sulphuric Acid Solution and 20 c c of Water. The leaves are packed into the percolator with a glass rod after the liquid has passed through, the flask is washed with 10 c c of the Chloroform-Ether mixture and the residue in the flask transferred to the percolator with several successive portions of 5 c c of the Chloroform-Ether mixture, and percolation continued with this menstruum, using in all 50 c c. The alkaloids are now removed from the Chloroform-Ether solution by shaking the separator vigorously, and the acid layer removed after complete separation. The Chloroform-Ether is shaken a second and a third time with 10 c c of a similar mixture of Sulphuric Acid and Water, the mixed acid liquids are transferred to a second separator, rendered distinctly alkaline with Ammonia Solution, and the liberated alkaloids shaken out first with 25 c c, then with 20 c c, and finally with 15 c c of Ether. The mixed ethereal solutions are evaporated on a water bath at a gentle heat, the residue dissolved in 3 c c of Ether and the Ether again evaporated. The residue is dissolved in 4 c c of Decinormal Volumetric Sulphuric Acid Solution, and the excess of acid is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Cochineal or Iodeosin Test Solution as an indicator of neutrality. If the number of c c of Fiftieth-normal Solution required be divided by 5, the quotient subtracted from 4, and the remainder multiplied first by 0.03 and then by 10, the product will represent the p c of Ether-soluble alkaloids present in the leaves.

The ash of Coca Leaves amounts to from 6 to 8 p c.

Preparation

EXTRACTUM COCÆ LIQUIDUM. LIQUID EXTRACT OF COCA
Percolate 20 of Coca Leaves in powder with Alcohol (60 p c)

until the drug is exhausted. Reserve the first 15 of percolate and evaporate remainder at a temperature below 80° C (176° F) to a soft extract, which dissolve in the reserved portion, and add Alcohol (60 p c), *q s* to yield 20 (1 in 1)

Note—As the Coca Leaves would be but imperfectly exhausted by the first 15 parts of the Alcohol, and as the active constituents are damaged or destroyed by heat, a fluid extract prepared by repercolation is much to be preferred. When thus prepared from carefully dried green leaves it contains 25 p c of solid Extract (dried at 212° F)

Dose.— $\frac{1}{2}$ to 1 fl ðim = 1 8 to 3 6 c c

Foreign Pharmacopœias—Official in Fr, Mex, Swiss and US. Not in the others

A dark green fluid extract prepared from Coca leaves. The *BP* product is not standardised, the *USP* preparation is required to contain 0 5 p c w/v of the Ether-soluble alkaloids of Coca, the *PG* does not contain a Fluid Extract

Tests—Fluid Extract of Coca possesses a specific gravity of 0 990 to 1 030, yields about 19 0 p c w/v of total solids, and contains about 50 p c w/v of Absolute Alcohol. The *BP* does not include a process for the determination of the alkaloids, the *USP* gives a method of determining the Ether-soluble alkaloids, the essential features of which are as follows.—A measured quantity of 10 c c of the fluid is introduced into a separator, rendered alkaline by the addition of 2 c c. of Ammonia Solution, and the alkaloids extracted by shaking for 1 minute with 25 c c of Ether. The separated aqueous liquid is shaken with a further quantity of 20 c c of Ether, the aqueous portion separated, the mixed ethereal solutions are shaken well for 1 minute with a mixture of 5 c c of Normal Volumetric Sulphuric Acid Solution and 5 c c of Distilled Water, the extraction being repeated with a mixture of 1 c c of Normal Volumetric Sulphuric Acid Solution and 9 c c of Water, the acid liquids in each case being separated and transferred to a second separator. Sufficient Ammonia Solution is now added to render the liquid distinctly alkaline, and the liberated alkaloids are shaken out with 20 c c of Ether, the extraction being completed with two further quantities each of 15 c c of Ether. The separated ethereal solutions are mixed, transferred to a 100 c c flask, the Ether evaporated and the residue dried. It is dissolved in 5 c c of Deci-normal Volumetric Sulphuric Acid Solution, and the excess of acid is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, using Cochineal or Iodocarmine Test Solution as an indicator of neutrality. If the number of c c of Fiftieth-normal Solution required be divided by 5, the quotient subtracted from 5, the difference multiplied first by 0 03 and then by 10, the product will represent the percentage w/v of the Ether-soluble alkaloids of Coca present in the sample. The average results of the examination by a special process, given in the reference, of 7 samples of Liquid Extract prepared by the official process and 7 'miscible' Liquid Extracts is recorded as 0 02, 1 421. The average yield of the 'official' samples was 0 38 p c, and they varied from 0 20 p c w/v to 0 816 p c w/v; the 'miscible' 0 091 p c of alkaloid, and they

valued from 0.014 to 0.294 p.c. w/v. A specimen of commercial 'miscible' Liquid Extract examined in the author's laboratory by the *U.S.P.* process showed 0.465 p.c. of alkaloids.

Not Official

EXTRACTUM COCÆ —A solid alcoholic green extract, prepared from carefully dried leaves.

Dose —2 to 10 grains = 0.13 to 0.65 gramme, in pills, pastils, or lozenges.

Foreign Pharmacopœias —Official in *Fr.* Codex, *Ital.* and *Span.* Not in the others.

ELIXIR COCÆ —Coca Leaves, 1, Simple Elixir, 6 —*Martindale*

Miscible Liquid Extract of Coca, 16.5, Simple Elixir to make 100 —*B.P.C.*

The Miscible Liquid Extract of Coca is 100 of *B.P.* Liquid Extract evaporated to 50, decanted from the deposit, which is washed with 10 of Water, the washings mixed with the concentrated fluid, and finally made up with Alcohol (60 p.c.) to 100.

TINCTURA COCÆ —Coca Leaves, 1, Alcohol (60 p.c.), 5 —*Fr.* and *Ital.*

Swiss, the same quantities, but with Alcohol (68 to 69 p.c.) All by weight.

This has been incorporated in the *B.P.C.* with Alcohol (60 p.c.) by measure.

VINUM COCÆ *Syn.* VIN DE COCA (*Fr.*) —Dried Leaves of Coca, 6, Vin de Malaga, 100, macerate for 10 days, and filter.

Wine of Coca can also be made by adding an equivalent quantity of the Liquid Extract to Wine.

Coca Wine, if sufficiently weak to be used as a beverage, requires a wine licence. The Excise has drawn the line at Wines containing $\frac{1}{2}$ grain of alkaloid to the oz., which would be at least twice as strong as the above.

Foreign Pharmacopœias —Official in *Fr.*, 6 in 100, *Mex.*, 3 in 100, *Span.*, 1 in 33 $\frac{1}{2}$, *Swiss.*, 1 of Fluid Extract in 20. *U.S.* about 1 in 15. Not in the others.

Elixir of Coca, 1, Detannated Sherry, to make 8 —*B.P.C.*

COCAINA.

COCAINE

METHYL-BENZOYL ECGONINE

$C_{17}H_{21}NO_4$, eq. 300.93

Large, colourless, odourless, monoclinic prisms, having a bitter taste, followed by anæsthesia of the mucous membrane. It is an alkaloid obtained from the leaves of *Erythroxylum Coca*, and its varieties.

Solubility —About 1 in 1300 of Water (Paul), 1 in 10 of Alcohol (90 p.c.), 1 in 50 of Olive Oil, 1 in 4 of Oleic Acid, 2 in 1 of Chloroform, 1 in 4 of Ether, 1 in 14 of Oil of Turpentine. Insoluble in Glycerin.

These figures have been incorporated in the *B.P.C.*

Official Preparation —Unguentum Cocainæ.

Not Official —Guttæ Cocainæ Oleosæ, Nebulæ Cocainæ Compositæ, Nebulæ Cocainæ Oleosæ, Oleatum Cocainæ, Unguentum Cocainæ, Unguentum Suprarenalin et Cocainæ.

Foreign Pharmacopœias —Official in *Fr.*, *Mex.*, *Span.* and *U.S.* Not in the others.

Tests—The *diagnostic* tests for Cocaine are the melting point, which should be 98°C ($208\ 4^{\circ}\text{F}$)—*B P* says 96° to 98°C (204.8° to $208\ 4^{\circ}\text{F}$), *USP* 98°C ($208\ 4^{\circ}\text{F}$), *Fr Codex* (1908) 98°C ($208\ 4^{\circ}\text{F}$)—the laevorotatory nature of its solutions, the alkaline reaction of its aqueous solution towards Litmus and Methyl Orange Solutions, the marked anæsthesia which it produces on the mucous membrane, and its mydriatic effect on the pupil of the eye. When exactly neutralised with Hydrochloric Acid the solution yields with Potassium Permanganate Solution a purple-violet precipitate, possessing a very characteristic microscopic appearance. This crystallisation takes place best in a solution of about 5 p.c. strength. When a crystal is moistened with one or two drops of fuming Nitric Acid, evaporated to dryness on a water-bath, and the residue moistened with a few drops of an alcoholic Potassium Hydroxide Solution, a peculiar characteristic fruity odour is produced. This test is characteristic of Cocaine, no other alkaloid extracted by Benzene from an ammoniacal solution behaving at all similarly. The salts of Cocaine are neutral to most indicators of neutrality, and the pure alkaloid may therefore be readily determined by titration with Normal or Deci-normal Volumetric Hydrochloric Acid Solution, 1 c.c. of Normal Acid representing 0.30093 grammes of the pure alkaloid. Iodocin or Cochineal Solution is the most suitable indicator for the purpose.

The impurities likely to be present in the alkaloid are also those more generally examined for in the Hydrochloride, and the methods by which they may be detected will be found under Cocaine Hydrochloride. Cocaine readily undergoes hydrolysis, and aqueous solutions are decomposed even on boiling, the decomposition being greatly facilitated by the presence of acid. The *B P* states, 'its solution in water acidulated with Hydrochloric Acid and the dry salt obtained on evaporating this solution afford the reactions mentioned under Cocaine Hydrochloride', the *B P C* has construed this expression into 'when its solution in Hydrochloric Acid is evaporated to dryness, the residue should respond to the tests given under Cocaine Hydrochloride', the *USP*, evidently with a view to minimising the amount of decomposition taking place, requires that it an alcoholic solution of Cocaine be carefully neutralised with Hydrochloric Acid and the solution evaporated to dryness, the residue should respond to the reactions and tests given under the Hydrochloride.

A solution of the alkaloid in water acidified with Nitric Acid should yield no opalescence or precipitate with either Silver Nitrate or Barium Chloride Solution. The alkaloid should yield no weighable residue on ignition. Only the Hydrochloride is official in the *L G*.

Preparation

UNGUENTUM COCAINÆ. COCAINE OINTMENT

Dissolve 1 of Cocaine in 4 (by weight) of Oleic Acid at a gentle heat, and mix with 20 of Lard. (1 in 25)

Oleatum Cocainæ—Cocaine, 5, Alcohol, 5, Oleic Acid, 50, Olive Oil, *q s* to produce 100—*U S*

Guttæ Cocainæ Oleosæ—Cocaine, 8 grains, Castor Oil, 1 oz—*St George's*

Not Official

NEBULA COCAINÆ COMPOSITA—Cocaine, 2 grains, Menthol, 4 grains, Eucalyptus Oil, 6 minims, Camphor, 4 grains, Spray Oil, 1 fl oz—*Bournemouth Formulary*

Nebula Eucalypti et Mentholis et Cocainæ, B P C, closely resembles the above—Cocaine, 0.5, Menthol, 1, Oil of Eucalyptus, 1.25, Camphor, 1, Liquid Paraffin, to produce 100

A modification of the above appears in the *B P C Supplement* as follows—**Nebula Cocainæ Composita**—Compound cocaine spray is prepared by dissolving 0.415 of Cocaine in sufficient compound Menthol and Thymol spray to produce 100

NEBULA COCAINÆ OLEOSA—Cocaine, 25 grains, Oil of Sweet Almonds, 1 fl oz Dissolve by heat—*Central Throat*

UNGUENTUM COCAINÆ—Cocaine, 2 grains, Soft Paraffin, 100 grains—*London Ophthalmic*

The title for this in *St Thomas's* and *B P C* is **Unguentum Cocainæ Dilutum** (pro oculis) The Cocaine should be finely powdered and rubbed with a small quantity of the Soft Paraffin, the remainder added, and dissolved with the aid of a gentle heat

Unguentum Cocainæ (Ophthalmic)—Cocaine Hydrochloride, 8 grains, Soft Paraffin, to 1 oz—*London*

UNGUENTUM ATROPINÆ ET COCAINÆ—*See Atropine*

UNGUENTUM SUPRARENALIN ET COCAINÆ—Suprarenalin, $\frac{1}{2}$ grain, Boric Acid, 1 grain, Cocaine Hydrochloride, 5 grains, Distilled Water, 15 minims, Hydrous Lanoline, 250 grains, Vaseline, 250 grains—*Bournemouth Formulary*

This has been incorporated in the *B P C* under the title **Unguentum Adreninæ et Cocainæ**, employing Adrenine

COCAINÆ HYDROCHLORIDUM.

COCAINE HYDROCHLORIDE

HYDROCHLORATE OF COCAINE—*B P '85*



FR CHLORHYDRATE DE COCAINE, GER, COCAINHYDROCHLORID, ITAL,
CLORIDATO DI COCAINA, SPAN, CLORURO DE COCAINA

Colourless, odourless, transparent, prismatic crystals, or acicular crystals, or a white glistening crystalline powder Taste slightly bitter, producing upon the tongue a tingling sensation followed by numbness of some minutes' duration

Solubility—2 in 1 of Water, 1 in $2\frac{1}{2}$ of Alcohol (90 p c), 1 in $2\frac{1}{2}$ of Glycerin, about 1 in 20 of Chloroform, almost insoluble in Ether, insoluble in fixed Oils

Medicinal Properties—Local anæsthetic, mydriatic Has been largely used for producing local anæsthesia in examinations of and operations on the eye and throat, and in dentistry ($\frac{1}{4}$ to $\frac{1}{2}$ grain

being injected into the gum), 2 p c solutions being used for the eye and 20 p c for the throat. It is used locally in producing anæsthesia of other mucous membranes, as the urethra, vagina, nose and rectum; in the form of spray containing $\frac{1}{2}$ to 2 p c, with or without other medicaments such as Adrenalin 1 in 5000, in aqueous solution, or $\frac{1}{2}$ p c of Cocaine with 1 p c of Menthol in Liquid Paraffin (see also p 772), in the form of Bougies, Pessaries, or Suppositories containing $\frac{1}{2}$ grain of the salt in each with Oil of Theobroma. Anæsthesia of the deeper seated tissues for minor operations is produced by local infiltration of Cocaine, combined generally, with Adrenalin. Injected locally for sciatica and for neuralgia. It has been used successfully as a preventive of seasickness, in doses of $\frac{1}{4}$ to 1 grain in solution, and in doses of $\frac{1}{8}$ grain every half-hour in the vomiting of pregnancy. As an ointment it is used in painful skin diseases, as shingles, in facial neuralgia and in pruritus.

Pastilles are made of various strengths from $\frac{1}{30}$ to $\frac{1}{8}$ grain in each, usually $\frac{1}{10}$ or $\frac{1}{15}$ grain. It is also supplied in granular effervescent form containing Cocaine Hydrochloride $\frac{1}{30}$, $\frac{1}{10}$, $\frac{1}{5}$, $\frac{1}{4}$ grain in each teaspoonful.

Hypodermic Tablets are supplied containing $\frac{1}{10}$, $\frac{1}{5}$, $\frac{1}{4}$, $\frac{1}{2}$ grain, also Cocaine Hydrochloride $\frac{1}{10}$ grain, Homatropine Hydrobromide $\frac{1}{10}$ grain, in each.

Hypodermic solutions are used containing 4 to 10 p c of the salt.

For external application in neuralgia, 10 or 20 p c solution of the alkaloid in Oil of Cloves, and a weaker solution 5 p c for toothache and earache. Cocaine applied on Lint or Cotton-Wool to a rigid os uteri is followed by m.

—*B M J* '98, ii 1874, '00, i 1340

In pertussis, dose $\frac{1}{10}$ grain three times daily for infants, increasing it according to the age, $\frac{1}{8}$ grain being given to children of 5 or 6 years —*L* '95, i 1429, *B M J E* '95, ii 28.

Combined with Opium in the internal treatment of cancerous disease —*B M J* '96, ii 718.

Four cases in which toxic symptoms have followed anæsthesia of the throat —*B M J E* '96, ii 95.

Used with a laryngeal syringe, shown (*B M J* '04, ii 1221) to be a satisfactory method of producing local anæsthesia during operation on the larynx.

Uncertain as a mydriatic, and cannot be relied upon to produce maximum dilatation of the pupil —*B M J* '99, ii 775.

As a method of the usual method of using Cocaine in operation upon the eye (*B M J* '04, ii 1303), an ointment has been advocated in combating photophobia (*B M J* '04, ii 1301), being stated not to produce desiccation of the corneal epithelium.

In cocainisation of the spinal canal, 2 c c of a freshly prepared sterilised 2 p c solution ($\frac{1}{2}$ grain), and the quantity should not be exceeded or toxic symptoms may result. Cocainisation following operation is checked by Phenacetin in 10-grain capsules or Hyoscine Hydrobromide —*L* '02, i 912, 1051.

$\frac{1}{2}$ -grain doses into spinal canal, preceded in some cases by hypodermic injection of 10 minims Liquor Strychninæ —*L* '02, ii 864.

Injection of Cocaine into the nerve-trunks about their point of division before an operation, and the administration of Morphine before it, tend to prevent shock (*L* '05, ii 579). For the induction of spinal anæsthesia in the treatment of Strychnine poisoning and of tetanus, 1 or 2 c c of a 1 p c solution of Cocaine Hydrochloride is injected into the spinal subdural space. Not more than $\frac{1}{8}$ grain should be injected at once, and it is well to commence with a much smaller quantity.

1 or 2 c c of a 3 p c solution of Eucaine B Hydrochloride may be used. The solution may be sterilised by boiling without loss of activity. Cocaine, and unpleasant or dangerous effects particularly have been used —*L* '05, ii 887.

Several fresh communications on the useful combination of Cocaine and Adrenalin have been made. The combined use is shown (*B M J E* '04, ii 60) to cause increase of the analgesic property and to lessen toxic effect. Solution recommended, 10 c c of a 1 in 200 Cocaine Hydrochloride Solution, 10 minims of a 1 in 1000 Adrenalin Solution. Their separate use has sometimes been advocated (*B M J* '04, ii 1227) in operation on the larynx, the Cocaine being used as a spray, a more diffuse effect being required. Adrenalin Solution has been used locally, its application being most desired where bleeding is taking place.

Suggested as probable that in the long run weaker solutions than 2 p c, say 1 in 100 to 1 in 400, will be used for intraspinal injection of Cocaine. The use of Eucaine suggested instead of Cocaine on account of its lesser toxicity and greater stability during sterilisation by heat—*L* '01, i 187.

Lumbar injections of 0.1 gramme (= $\frac{1}{10}$ grain) during labour—*L* '01, ii 365, 645.

The physiological effects of cocainisation of the spinal canal—*L* '01, ii 1280.

Dangers of anaesthesia by injection of Cocaine into the spinal canal—*L* '01, i 975.

An objection to the lumbar method of producing anaesthesia being employed as a routine practice on a large scale is the poisonous and treacherous character of the drug hitherto used, viz, Cocaine—*B M J* '07, ii 869.

For local anaesthesia in the extraction of teeth, the best results are obtained with 1 p c solution of Cocaine combined with 5 p c of Adrenalin Chloride. Danger of syncope and other accidents from Cocaine, due in most cases to carelessness or ignorance in the method, the same amount of Cocaine is more dangerous in a concentrated than in a weaker solution—*B M J* '07, i 895.

Use of 10 p c solution for extraction of teeth strongly deprecated, 1 p c solution perfectly effective, dose should never exceed 1 grain—*B M J* '07, i 788, 848. Death from urethral injection of 3 grains—*B M J* '06, ii 868.

Cocaine intoxication and its demoralising effects—*B M J* '02, i 1020, 1041.

Dose— $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.01 to 0.03 gramme.

Ph Ger maximum single dose, 0.05 gramme, maximum daily dose, 0.15 gramme.

Prescribing Notes—*Unless a preservative be used, solutions should be freshly prepared to prevent the development of a fungus. As solutions of Cocaine are damaged by heat, they must not be sterilised by boiling. Salicylic Acid is the best, if not the only effectual preservative for aqueous solutions of Cocaine, but it is very irritating to the eye. As Borax is incompatible with this salt, an equivalent quantity of Boric Acid should be prescribed.*

Incompatibles—Alkalis and alkaline Carbonates, Borax, Carbolic Acid, Mercurous and Mercuric Chlorides, and the majority of soluble Silver salts.

Official Preparations—Injectio Cocainæ Hypodermica, and Lamellæ Cocainæ. Used in the preparation of Trochiscus Krameriae et Cocainæ.

Not Official—Guttæ Cocainæ Hydrochloridi, Nebula Cocainæ, Pastillus Cocainæ, Pastillus Cocainæ et Morphinae, Trochisci Cocainæ, Trochisci Cocainæ et Morphinae, Cocainæ Citras, Cocainæ Hydrobromidum, Cocainæ Lactas, Cocainæ Nitrates, Cocainæ Oleas, Cocainæ Phenylas, Cocainæ Salicylas, Cocainæ Sulphas, Eucaine, Eucaine Hydrochloride, Eucaine Lactate, Orthoform, Orthoform Hydrochloride, Benzoyl-Pseudotropine, Holocaine, Holocaine Hydrochloride, Acoine Nirvanin, Nervocidine, Alypin, Novocaine, Stovaine.

Antidotes—Inhalation of Nitrite of Amyl—*B M J* '87, i 625, 695, 1401, 88, i 757. Strychnine and Digitalin—*L* '98, i 718.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Russ., Span., Swed., Swiss and U.S. Not in the others.

Tests—Cocaine Hydrochloride possesses a melting point according to the *B.P.* of from 180° to 186° C (356° to 366.8° F), the *U.S.P.* states 189° C (372.2° F), mentioning that the presence of minute quantities of impurities may reduce the melting point to

180° C. (356° F), or even less, the *Fr Codex* (1908) gives 186° C (366 8° F), the *PG* gives 183° C (361 4° F) It should be perfectly free from colour, should be readily and completely soluble 2 in 1 of Water, forming a perfectly colourless solution, which should be absolutely neutral to Litmus paper Its solutions are laevogyrate Its aqueous solution yields with Ammonium Carbonate Solution and with Potassium, Sodium, or Ammonium Hydroxide Solution a white precipitate, amorphous when precipitated from strong solutions, but rapidly becoming crystalline If this precipitate be dissolved in Ether, the ethereal solution separated, and the Ether carefully evaporated, the residue should respond to the distinctive tests given in the large type under 'Cocaine' The salt should dissolve without change of colour in pure concentrated Sulphuric Acid or in pure Nitric Acid On warming its solution in the former acid, it chars, at the same time evolving an agreeable aromatic odour, and yielding a crystalline sublimate of Benzoic Acid Its aqueous solution affords, with Potassio-mercuric Iodide (Mayer's) Solution, a white precipitate, precipitation occurring even in very dilute solution, with Auric Chloride Solution, a yellow precipitate, with Sodium Bichlorate Solution, a white precipitate, with Picric Acid Solution, a yellow precipitate, rapidly becoming crystalline, with Mercuric Chloride Test Solution, slightly acidulated with Hydrochloric Acid, a white precipitate soluble in hot Water, with Platinic Chloride Solution, a yellow crystalline precipitate, with Palladous Chloride Solution (5 p c), followed by the addition of Chlorine Water, a red precipitate, with Chromic Acid Solution or with Potassium Bichromate Solution, followed by the addition of Hydrochloric Acid, a yellow crystalline precipitate An aqueous solution of the salt should yield with Silver Nitrate Solution a curdy white precipitate, insoluble in Nitric Acid, and which, when filtered and washed is readily soluble in Ammonia Solution or in Potassium Cyanide Solution

A mixture of equal parts of the salt and Mercuric Chloride is blackened when moistened with diluted Alcohol

Iodine or Iodo-potassium Iodide (Wagner's) Solution precipitates Cocaine from its aqueous solutions, in very dilute solution the precipitate appears of a rose colour, in stronger solutions, brown Upon the reaction of Cocaine with Iodine solution has been founded (*PJ* '01, 1553, '01, 222, *JSCI* '01, 1031, *JCS Trans*, '01, 675) a process for the determination of Cocaine The Cocaine solution should contain about 1 gramme of the alkaloid in the form of a salt in 100 cc The Deci-normal Volumetric Iodine Solution should be added in excess, and the excess of Volumetric Iodine Solution titrated with Deci-normal Volumetric Sodium Thio-sulphate Solution Cocaine can be fairly accurately determined by this method, in the presence of Ecgonine, but not in the presence of Benzoyl Ecgonine As, however, neither Ecgonine nor Benzoyl Ecgonine are extracted from aqueous alkaline solution by Petroleum Ether or Ether, a method of separation is available Cocaine is readily extracted by these immiscible solvents Assuming the production of Cocaine Di-iodo-hydriodide as a result of the reaction, one

molecular quantity of Cocaine is necessary for each molecular quantity of Iodine used

The more generally occurring impurities are Cinnamyl-Cocaine, Cocamine and other products derived from Cocaine, amorphous alkaloids, Sulphates, excess of moisture, and mineral matter. A great deal of controversy has raged round the tests adopted for the detection of these impurities. As a test for the presence of Cinnamyl-Cocaine, Cocamine and other products derived from Cocaine, the *BP* utilises the Permanganate Test. An excess of Potassium Permanganate Solution is added to a 1 p c solution of the Cocaine salt, when a copious red precipitate is produced, which is required not to undergo any alteration in colour within an hour. It is not clear how the *BP* proposes to observe the alteration in colour of the red crystalline precipitate in the presence of an excess of Potassium Permanganate Solution. The value of results yielded by the Potassium Permanganate Test depends entirely upon the method of carrying it out, and the conditions appear to have been completely misinterpreted, the test as officially described is therefore worthless. The Cocaine solution requires to be of such a dilution that Cocaine Permanganate is not precipitated, and the Potassium Permanganate Solution requires to be sufficiently weak to just colour the liquid. The test may be well applied as follows — Dissolve 0.1 gramme of the salt in 5 c c of Water acidulated with 3 drops of Diluted Sulphuric Acid and add 0.5 c c of a 1 in 1000 Potassium Permanganate Solution. The colour should not disappear within an hour. The *USP* test is on similar lines to the above, but 3 drops of Deci-normal Volumetric Potassium Permanganate Solution are used, and the violet colour which is produced should not fade in half an hour. The *PG* uses 5 drops of Potassium Permanganate Solution (0.1 p c w/w), and states that when protected from dust no perceptible decolorisation should occur in half an hour.

According to Paul and Cownley, the most efficient test for ascertaining the purity of the salt for medicinal purposes is MacLagan's Test. 1 grain of the salt is dissolved in 2 fl oz of Water, 3 drops of Ammonia Solution (*BP*) added, and the mixture stirred briskly with a glass rod, within a few minutes a crystalline precipitate should be thrown down, leaving no turbidity in the supernatant liquid. The *BP* idea of the test is as follows — A weighed quantity of 0.1 gramme of the salt is dissolved in 100 c c of Water and 0.25 c c of Ammonia Solution added. It should afford a clear solution from which a crystalline deposit should gradually separate on stirring. The quantities given in the *BP* translated into the terms of MacLagan's Test indicate 0.86 grain of the salt in 2 fl oz of Water and 5 drops of Ammonia Solution, or 1 grain of the salt in a little over 2½ oz of Water and 5 drops of Ammonia Solution, it will therefore be observed that the proportions are not strictly maintained. The *USP* maintains the same relative proportions of Cocaine salt to Water as in MacLagan's Test, but uses 4 instead of 3 drops of Ammonia Solution, and specifies 15 minutes as a limit of time for the appearance of the crystalline precipitate, the mixture

should be vigorously stirred, and the sides of the beaker rubbed occasionally with the stirring rod. The *PG* adopts the same proportions of salt and Water as the *BP*, but after the addition of the Ammonia Solution allows the liquid to remain at rest for an hour, when no opalescence should be produced. The essential feature of the test is the strict maintenance of the original proportions.

Maclagan's Test has been subjected to a good deal of adverse criticism on the continent. Gunther (*CD* '99, i 457) claimed to have discovered a new base (Ethyl-benzoyl-ecgonine), possessing a melting point of 110° to 111° C (230° to 231° F), and to have shown that the salts of Cocaine as generally found on the market, and as hitherto obtained, are not solely a compound of the alkaloid Cocaine with an individual acid, but that the Cocaine is associated with an isomeric alkaloid. The isomeric Hydrochloride gives the Maclagan's Ammonia reaction quickly and distinctly even in 1 in 2500 solution, whereas a 1 in 1000 Cocaine Hydrochloride solution does not give any crystallisation whatever on the addition of Ammonia Solution as prescribed by the test. He concludes, therefore, that the crystallisation is not due to the Cocaine at all, but to the isomeride. Although endeavours have been made (*PJ* '99, i 25) in various ways to obtain evidence of the existence of this high melting point base, no base having a melting point of 111° C (231° F) could be obtained, and only 0.00006 of a base melting at 104° to 108° C (219° to 222° F), which was quite insufficient to account for the crystalline precipitate in Maclagan's Test.

Notwithstanding the adverse criticism to which the Maclagan's Test has been subjected, it has been conclusively shown (*PJ* '98, i 449, 473, 586, '99, i 431, 523, 524, '99, ii 25, 66, *CD* '98, i 511, '99, i 897, *Merck's Annual Report*, '99, 51) that it affords the best guarantee of the purity of Cocaine Hydrochloride, and that any sample which does not satisfy the test should not be regarded as sufficiently pure for pharmaceutical purposes.

Schaefer (*CD* '99, i 591, *PJ* '99, i 336) formulates a new test depending upon the relative solubilities of Cocaine Chloride and the residual alkaloidal Chlorides in Water and in Water acidulated with Hydrochloric Acid. A weighed quantity of 0.05 gramme of the specimen is dissolved in 20 c.c. of Water, mixed with 5 c.c. of a 3 p.c. Chromic Acid solution, and 5 c.c. of a 10 p.c. Hydrochloric Acid Solution is added, the temperature being maintained at 15° C (59° F). If more than traces of foreign Coca-bases be present, the solution becomes at once cloudy, if the Cocaine Hydrochloride be pure, a clear solution will result.

A new alkaloid isomeric with Cocaine, and called by Schaefer, Cocainidine, is stated (*CD* '99, i 602, *PJ* '99, i 359) to have been found in Coca Leaves, and in commercial samples of Cocaine Hydrochloride. Schaefer's Chromic Acid Test has been critically examined in the author's laboratory (*CD* '99, i 641, 702). Six specimens of pure Cocaine Hydrochloride obtained from manufacturers of the highest repute were subjected to the test. All gave distinct turbidity, either immediately or after a few seconds. A specimen of Cocainidine

Hydrochloride was also examined, and gave a distinct turbidity at once. Dr Schaefer's Chromic Acid Test therefore produces a turbidity in solutions of the best commercial samples, and also of his Cocamidine Hydrochloride. The base obtained by treating an aqueous solution of Cocamidine Hydrochloride with Ammonia Solution, when washed, and dried over Sulphuric Acid, had the same melting point as the base obtained on similarly treating one of the best samples of Cocaine Hydrochloride, viz., 98°C (208.4°F).

It was subsequently mentioned (*CD* '99, 1 745) that the specimen of Cocamidine Hydrochloride submitted contained Isotropyl-Cocaine.

It has been demonstrated (*CD* '99, 1 897, *PJ* '99, 1 523) that the concentration of the Hydrochloric Acid plays an important part in the test, as does also the age of the Chromic Acid solution. The same sample of Hydrochloride gave a precipitate on the addition of 10 p c acid, but with 7.5 p c the solution remained clear, whilst another sample was satisfactory with 10 p c of acid, but precipitated with 12.5 p c. A freshly-prepared Chromic Acid solution, which remains quite clear when added to solution of the sample, after 24 to 48 hours' standing will produce a precipitate.

Schaefer endeavours (*PJ* '99, 1 66) to meet the criticisms which have been levelled against the test by stating that the finest commercial brands of Cocaine Hydrochloride answer the test satisfactorily. The Chromates of the amorphous alkaloids are far less soluble in solutions containing the higher percentage of Hydrochloric Acid. An acid of 10 p c was decided upon, and if the reaction is carried out with a stronger acid it is necessary to do so side by side with a specimen of chemically pure Cocaine, as at a low temperature Cocaine Chromate causes a turbidity in the more acid solution. His own experience fails to confirm Merck's remarks *re* the age of the Chromic Acid solution. The all-important feature of the test is that a temperature of 15°C (59°F) should be maintained. Notwithstanding this the value of the Chromate Test is generally held to be questionable, and there is no doubt that the MacLagan Test is far preferable. Cownley (*PJ* '99, 1 336, '99, 11 66) is of opinion that probably the only salt that would pass the test would be one prepared from synthetic Cocaine.

Both *USP* and *PG* includes a test with Chromic Acid Solution. The *USP* employs 5 c c of a 2 p c solution of the Cocaine salt, the *PG* a solution of 0.5 gramme of the alkaloidal Hydrochloride in 5 c c of Water, and both add 5 drops of a 5 p c Chromium Trioxide Solution. A yellow precipitate is produced, which redissolves on shaking the mixture, but is reprecipitated as a permanent orange-coloured crystalline precipitate on the addition of 1 c c of Hydrochloric Acid.

Messrs Zimmer and Co point out (*PJ* '99, 11 315) that a salt which does not give a crystalline precipitate by MacLagan's Test within 5 minutes should be rejected as being too impure, since the time required to obtain a precipitate serves as an index of purity.

The aqueous solution of Cocaine Hydrochloride when acidified with Diluted Hydrochloric Acid shall yield no opalescence or

precipitate with Barium Chloride Solution, indicating the absence of Sulphates

The formula given in the *B P* represents the anhydrous salt. It is officially required to lose not more than 1.0 p.c. of moisture as determined by drying a weighed quantity for 20 minutes at a temperature of 95° to 100° C (204° to 212° F), the *P G* states that it suffers no loss of weight at 100° C (212° F). The Brussels Convention recommends only the anhydrous salt should be recognised. The absence of mineral matter is at once detected by the residue left on a glass plate; no weighable residue should remain when 0.5 to 1 gramme of the salt is heated to redness with free access of air.

Preparations

INJECTIO COCAINÆ HYPODERMICA. HYPODERMIC INJECTION OF COCAINE

Dissolve $\frac{1}{2}$ gram of Salicylic Acid in 6 fl drms of boiled Distilled Water, add 33 grains of Cocaine Hydrochloride, dissolve, and if necessary add Distilled Water (recently boiled and cooled), *qs* to produce 6 fl drms (1 in 10)

Dose—By subcutaneous injection, 2 to 5 minims = 0.12 to 0.3 cc

11 minims contain 1 grain of Cocaine Hydrochloride

Foreign Pharmacopœias—Official in Mex and Span 1 in 100

LAMELLÆ COCAINÆ. DISCS OF COCAINE

Gelatin discs, containing $\frac{1}{100}$ grain of Cocaine Hydrochloride

Four times the strength of *B P* '85, which contained $\frac{1}{400}$ grain

Ophthalmic discs, each containing about 0.0005 gramme = $\frac{1}{2000}$ grain Cocaine Hydrochloride, are official in Ital Ph

They are also supplied containing Cocaine $\frac{1}{100}$ grain with Atropine $\frac{1}{1000}$ grain, Cocaine $\frac{1}{100}$ grain with Physostigmine $\frac{1}{1000}$ grain, and Cocaine $\frac{1}{100}$ grain with Homatropine $\frac{1}{100}$ grain

TROCHISCUS KRAMERIÆ ET COCAINÆ. See KRAMERIA

Not Official.

GUTTÆ COCAINÆ HYDROCHLORIDI—Cocaine Hydrochloride, 10 grains, Distilled Water, 1 fl oz—*London Ophthalmic*

NEBULA COCAINÆ—Cocaine Hydrochloride, 48 grains, saturated solution of Boracic Acid, 1 fl oz—*Central Throat*

PASTILLUS COCAINÆ— $\frac{1}{10}$ grain of Cocaine Hydrochloride in each (Throat), $\frac{1}{20}$ grain (*St Thomas's*). *Fr* has Tablettes de Chlorhydrate de Cocaine, each containing $\frac{1}{4}$ grain of Cocaine Hydrochloride

PASTILLUS COCAINÆ ET MORPHINÆ—Cocaine $\frac{1}{10}$ grain, Morphine Hydrochloride, $\frac{1}{30}$ grain—*Martindale*, incorporated in the *B P C*

Useful for coughs

TROCHISCI COCAINÆ.— $\frac{1}{10}$ grain—*Central Throat and Throat*

TROCHISCUS COCAINÆ ET MORPHINÆ—Cocaine Hydrochloride, $\frac{1}{20}$ grain, Morphine Hydrochloride $\frac{1}{40}$ grain—*St George's*

COCAINÆ CITRAS—Colourless, hygroscopic crystals, readily soluble in Water Used in dentistry

COCAINÆ HYDROBROMIDUM—Transparent prisms, soluble in Water

COCAINÆ LACTAS—A white semi solid, readily soluble in Water
Used as an injection in tubercular cystitis

COCAINÆ NITRAS—Large, tabular crystals, readily soluble in Water and in Alcohol (90 p c) Used in ophthalmic practice and in urethral surgery in conjunction with Silver Nitrate

Tests.—The aqueous solution yields the tests distinctive of Cocaine given under Cocaine and Cocaine Hydrochloride The aqueous solution decolorises Indigo Sulphate Solution containing an excess of Sulphuric Acid When a solution of Ferrous Sulphate is poured carefully upon a well-cooled mixture of equal parts of Sulphuric Acid and a solution of the salt, a brown or purple brown coloration is formed at the junction of the two fluids When warmed with Copper and Sulphuric Acid reddish-brown fumes are evolved It should leave no weighable residue when ignited with free access of air

COCAINÆ OLEAS—A crystallisable salt, insoluble in Water, soluble in Oleic Acid and fixed Oils Is useful when a salt of Cocaine is required in a fatty basis such as an ointment or suppository

COCAINÆ PHENYLAS (Cocaine Carbolate)—A yellow, or yellowish-brown, semi solid mass Insoluble in Water, soluble in Alcohol (90 p c) and in Ether Introduced as a local anæsthetic, analgesic and sedative Combined with Acetanilide it is stated to have been found useful in gastralgia in doses of $1\frac{1}{2}$ grains daily 1 drop of a 10 p c alcoholic solution of the salt has been found useful in conjunctival catarrh Has also been found useful in dentistry Owing to its insolubility is not so rapidly absorbed, and the action is more prolonged

Dose—Internally, $\frac{1}{12}$ to $\frac{1}{8}$ grain = 0.005 to 0.01 gramme

COCAINÆ SALICYLAS—Short, thick, somewhat deliquescent crystals, soluble 5 in 1 of Water, $2\frac{1}{2}$ in 1 of Alcohol (90 p c)

Dose— $\frac{1}{8}$ to $\frac{1}{2}$ grain = 0.01 to 0.032 gramme

Has been recommended hypodermically in asthma

Tests—Cocaine Salicylate dissolves readily in Water, the aqueous solution yields the reactions distinctive of Cocaine given under Cocaine and Cocaine Hydrochloride

The diluted aqueous solution yields with Ferric Chloride Test-solution a violet coloration

COCAINÆ SULPHAS—In prisms, or as a white, granular powder, soluble in Water

EUCAINÆ—Under this name two basic principles have been introduced
Eucaine (A) (Methylester of Benzoyl *n* methyl tetra methyl-gamma oxy piperidine carboxylic Acid) **Eucaine (B)** (Benzoyl vinyl di aceton alkaline) Synthetic products resembling Cocaine both in chemical and physiological action

Eucaine (B) is the base in general use It is insoluble in Water, but readily soluble in Alcohol (90 p c) and in Ether Soluble 1 in $1\frac{1}{2}$ of Aniline Oil

EUCAINÆ HYDROCHLORIDE—There are two salts bearing this name, marked A and B, and prepared from the corresponding bases Eucaine A and Eucaine B The B salt is that generally used in medicine, and it is the practice to dispense it when Eucaine Hydrochloride is ordered unless the A is specified.

Eucaine B Hydrochloride ($C_{15}H_{21}NO \cdot HCl$, eq 281.54)—A fine white odourless crystalline powder, possessing a bitter taste followed by a feeling of numbness of the tongue

Solubility—1 in 40 of Water, 1 in 12 of Alcohol (90 p c), 1 in 4 of Aniline Oil

Medicinal Properties—A powerful local anæsthetic It is not so generally effective as Cocaine, but is less toxic Solutions of Eucaine salts may be sterilised by boiling without undergoing decomposition

The B salt is superior to Eucaine Hydrochloride (A) for use in ophthalmic work, as it is free from the irritating effects of the latter, and is an equally powerful local anæsthetic. Used in 2 p.c. aqueous solution 2 drops applied every three minutes until 10 drops have been used.—*B M J* '97, i 184, '97, ii 1560, *L* '00, i 1106

A 2 p.c. solution recommended for hypodermic use, 40 minims in two doses of 20 minims each distributed over three or four places.—*L* '99, ii 552

5 to 8 p.c. usual strength of solution necessary to produce anæsthesia for minor surgical operations. Usual dose 20 minims, and as much as 120 minims used without unfavourable results.—*L* '99, i 137, '99, ii 318

10 to 15 minims of an approximately 5 p.c. solution made with equal parts of Aniline Oil and Alcohol (90 p.c.) for the production of local anæsthesia of the ear.—*L* '00, i 1125

Solution recommended by Baker.—Eucaine Hydrochloride B, 1, Sodium Chloride, 8, Sterilised Water, 1000, Injection of 10 c.c. As much as 10½ oz injected without any ill effects. Such an amount, however, rarely necessary. Powders containing sufficient of the two salts for two ordinary operations may be kept ready, and can be dissolved in the necessary quantity of water and boiled before use.—*L* '99, i 282, '00, i 156

Addition of 0.8 p.c. Sodium Chloride to solution of Eucaine improves its analgesic properties, and has also other desirable effects

An improved solution (Baker's) for the production of local anæsthesia.—B Eucaine Hydrochloride, 0.2 gramme, Sodium Chloride, 0.8 gramme, Adrenalin Chloride, 0.001 gramme, Distilled Water, 100 grammes.—*L* '03, ii 204

Hermiotomy performed under local anæsthesia produced by the injection of 40 minims of a 1 p.c. solution, followed by 20 minims more of the same solution dropped into the wound during operation.—*L* '03, ii 580

Hypodermic injection of 40 minims of a 7½ p.c. solution Eucaine Hydrochloride (equal to 2½ grains of the drug) before operation of suprapubic cystotomy.—*L* '00, i 928

The use of a solution of Cocaine Hydrochloride 10, B Eucaine 10, Aniline 50, Alcohol (90 p.c.) 50, for the production of local anæsthesia in the ear, nose and throat, in order to obviate the dangers of using strong Cocaine solutions. To avoid change of colour, the solutions are best kept separately, e.g., as a 20 p.c. solution of Cocaine in Alcohol (90 p.c.), and a 15 to 20 p.c. solution of Eucaine (B) in Aniline Oil.—*L* '01, i 698

30 minims of a 5 p.c. solution injected around bed of the finger-nail causes swelling and unhealthy blueness of the skin of finger tip.—*L* '01, i 1510. Pointed out in reference to above that Eucaine solutions should be boiled immediately before use, injected at body temperature, and the use of a syringe previously used for Morphine should be avoided.—*L* '01, i 1648

Solutions recommended in ophthalmic work, 2 p.c., in the urethra and bladder, 2 p.c., for the nose and throat or as a paint or spray for mucous surfaces generally, 5 to 10 p.c., and in dental work, 2 to 5 p.c.—*B M J E* '03, i 36

The relative toxicity of Cocaine and Eucaine.—*T G* '99, 689

With reference to the use of strong solutions of these salts for hypodermic injections, it is pointed out they should not be used of greater strength than 3.0 p.c., stronger solutions, being hypotonic with the blood, are liable to cause sloughing of the tissues. Much useful information on the subject is given in a paper in the *B M J* '04, ii 1862, and the formula for a suitable solution is there given as Beta-Eucaine Hydrochloride, 0.2 gramme, Sodium Chloride, 0.8 gramme, Adrenalin Chloride solution (1 in 1000), 10 minims, Distilled Water, sufficient to make 100 c.c.

On the pharmaceutical side, the incompatibility of Salicylic Acid with Eucaine may be drawn attention to. This is a point of some importance, as the Pharmacopœia directs Salicylic Acid to be used as a preservative for Injectio Cocaine Hypod., but an attempt to carry this practice out with the Eucaine salt will result in precipitation, as the Salicylate is much less soluble.

Spinal anæsthesia induced by Eucaine B in the treatment of tetanus.—*L* '05, ii 888. The solution used consisted of Eucaine B Hydrochloride, 1½ grains, Morphine Sulphate, ½ grain, Sodium Chloride, 8 grains, Water, to

3½ oz From 15 to 16 c c of cerebrospinal fluid were withdrawn, and injections of from 3 to 4 c c of the above fluid given at various intervals

In sciatica (*B M J E* '05, i 44), 72 to 100 c c of 10 p c solution of Eucaine in saline solution are injected into the nerve at its point of emergence

It is by no means uncommon to be asked to make up solutions containing as much as 4 or 5 p c Although this can be accomplished easily by the aid of heat, the salt does not remain in solution, and even 3.2 p c solutions when prepared by the aid of heat do not remain bright long even when kept in hermetically sealed capsules, but soon deposit tufts of crystals On inquiry of the manufacturers, a letter resulted acknowledging the correctness of the *Companion* figures, and pointing out that as the solution has to be used at the body temperature, any salt which crystallises out will again be taken into solution The use of the stronger solutions of Eucaine has, however lately been deprecated, and the valuable work recorded in the *B M J* has shown that inasmuch as they are hypotonic with the blood, they are liable to produce necrosis of the tissues if used for the purposes of hypodermic injection

Foreign Pharmacopœias—Official in Dan and Swiss (Trimethylbenzoxypiperidinium-hydrochloricum)

Tests—Eucaine (B) Hydrochloride possesses a melting point of 268° C (514.4° F) A weighed quantity of 0.1 gramme dissolved in 10 c c of Water yields on the addition of 1 drop of Ammonia Solution a crystalline precipitate, which redissolves, but is again thrown down on the further addition of Ammonia The precipitate caused by 4 drops of Ammonia Solution completely dissolves on the further addition of 20 c c of Water On again adding 4 c c of Ammonia Solution the precipitate forms again, and is again dissolved on the addition of 10 c c of Water, but no further precipitate is formed on the addition of Ammonia Solution, only a milky cloudiness, disappearing on the addition of Water

A 1 p c aqueous solution yields with Potassium or Sodium Hydroxide Solution a precipitate of the free base, which dissolves readily in Ether Mercuric Chloride Solution produces no precipitate The aqueous solution yields a precipitate with Mayer's reagent and lemon yellow precipitate with Picric Acid Solution If a crystal be moistened with Nitric Acid and evaporated to dryness, it leaves a colourless residue, which evolves a characteristic odour of Benzoic Acid Ethyl-Ester when moistened with an alcoholic Potassium Hydroxide Solution

The aqueous solution acidified with diluted Nitric Acid yields on the addition of Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, readily soluble in Ammonia Solution

It may be distinguished from Alpha-Eucaine by the 1 p c aqueous solution yielding no precipitate on the addition of a few drops of 10 p c Potassium Iodide Solution, Alpha-Eucaine Solution yielding under similar conditions a crystalline precipitate

It may be distinguished from Cocaine by mixing a little of the salt with some Mercurous Chloride and moistening the mixture with Alcohol (90 p c), when no darkening in colour should be noticed Cocaine under similar conditions immediately darkens in colour Eucaine (B) Hydrochloride should dissolve without change of colour in concentrated Sulphuric or Nitric Acid

It should leave no weighable residue upon ignition with free access of air

Eucaine Lactate ($C_9H_{11}NO_5$, eq 334.72)—A white odourless crystalline powder possessing a bitter taste, and subsequently producing a feeling of numbness of the tongue It has been introduced as a local anæsthetic, which is claimed to be more soluble than the Hydrochloride

Solubility—1 in 4 of Water, 1 in 8 of Alcohol (90 p c)

It is usually employed in the form of a 2 to 3 p c solution

ORTHOFORM—This base is Para-amido-meta-hydroxybenzoic Acid Methyl Ester, a synthetic product introduced as a substitute for Cocaine A white, odourless, tasteless, crystalline powder, or in colourless crystals melting at 120° C Soluble, 1 in 450 of Water, 1 in 6 of Alcohol (90 p c), 1 in 12½ of Ether

Another base, Meta-amido para-hydroxybenzoic Acid Methyl Ester, has been introduced under the name of 'Orthoform New' Solubility is practically the same as above

Medicinal Properties.—Local anæsthetic employed in ulcerations of the upper air passages Useful where nerve endings are exposed, but has no action on unbroken skin and but little on healthy mucous membrane Best administered as a spray, using 10 p c solution made with Alcohol (45 p c), but the powder may be employed either alone or mixed with an equal quantity of Lycopersodium for insufflation, or in the form of a 10 p c ointment, a saturated solution of Orthoform in Collodion is used as a varnish Said to be of value as an anodyne in ulcer or cancer of the stomach in doses of 8 to 16 grains An aqueous solution of the Hydrochloride is used as a paint —*B M J* '98, 1 362, *Pr* lx1 505

Non-toxic and powerfully antiseptic On account of its sparing solubility it is but slowly absorbed Nearly 2 oz have been employed in the course of a week for dusting wounded surfaces without injurious effect —*B M J E*, '97, 11 55, *P J* '97, 11 277, *B M J* '98, 1 362

As an ointment it is useful in burns, in ulcers of the leg and in syphilitic ulcers —*L* '98, 1 1024, *B M J E* '98, 1 76, *P J* '98, 11 661

Used (suspended in Glycerin) for intra-uterine medication —*L* '98, 1 1434

Cotton-Wool plug steeped in an alcoholic solution introduced into the cavity of a tooth for the relief of toothache —*T G* '99, 270, *P J* '99, 1 83

In fissure of the nipple —*T G* '99, 337

As an insufflation in stomatitis in children —*B M J E* '99, 1 75

Insufflated, dusted on, or used as an ointment is most efficient (*B M J* '05, 11, 1008) in allaying pain, in burns, ulcerative stomatitis, tuberculosis and malignant ulceration, whether of the larynx or other regions Or it may be given internally up to 8 grains for gastric ulcer, carcinoma, or nervous dyspepsia

As an emulsion, Orthoform, 25, Olive Oil, 100, as an insufflation, 10 to 20 c mgrains or as a 10 p c aqueous solution of the Hydrochloride for laryngeal application —*B M J E* '99, 1 20, 64

ORTHOFORM HYDROCHLORIDE —A white, crystalline powder, which is soluble, 1 in 8 4 of Water, 1 in 17 of Alcohol (90 p c) Insoluble in Ether It may be employed for internal administration or for urethral injection, but is too acid for hypodermic injection or application to the eye —*L* '97, 11 733, *B M J E* '97, 11 55 Injection of a 10 p c solution in gleet —*L* '97, 11 738

Dose.—1 to 5 grains = 0 06 to 0 32 gramme

BENZOYL-PSEUDOTROPEINE (Tropacocaine, Tropain) —First obtained from Java Coca Leaves and afterwards made synthetically The Hydrochloride has been used to produce anæsthesia of the eye during operations, it is much less toxic than Cocaine —*B M J* '92, 11 406, '94, 11 598, *L* '94, 11 598, *T G* '94, 653, *MA* '93, 52

0 05 gramme (= $\frac{1}{4}$ grain) in 1 c c (16 minims) Water as an injection into the spinal canal to produce analgesia —*B M J E* '02, 1 75

0 05 gramme (= $\frac{1}{4}$ grain) dissolved in 5 c c (80 minims) of cerebrospinal fluid and reinjected to induce anæsthesia without undesirable concomitants —*Merck's Report*, '02, 166

Intraspinal injection of 1 c c of a 5 p c solution in puerperal eclampsia —*B M J E* '02, 11 6

Intraspinal injection in doses of 0 07 gramme ($1\frac{1}{4}$ grain) —*L* '05, 11 561

Of the drugs that now hold the field in lumbar anæsthesia, Stovaine, Novocaine, Alypin, and Tropacocaine, the most recent publications indicate a growing preference for the last named as the most reliable and the least dangerous —*B M J* '07, 11 1002, and 873

For the treatment of 1 c c of a 5 p c solution, addition of Adrenalin wholly unnecessary —*B M J* '07, 11 873

HOLOCAINE (Para-diethoxy-ethenyl-diphenyl-amidine) —A synthetic product introduced as a substitute for Cocaine In colourless crystals which melt at 121° C (249 8° F) Insoluble in Water, readily soluble in Alcohol (90 p c) and Ether

A powerful base, forming sparingly soluble salts with acids

HOLOCAINE HYDROCHLORIDE —The Hydrochloride of the above base. Occurs in colourless crystals, soluble in Water, Alcohol, and Ether

Solubility—1 in 50 of Water, 1 in 6 of Alcohol (90 p c)

Medicinal Properties—Used in the form of 1 p c solution in ophthalmic surgery. Produces complete and rapid anaesthesia without pain, and neither dilates the pupil nor affects the blood-vessels. On account of its toxicity, it cannot be used hypodermically. Its instillation into the eye causes a slight feeling of burning which rapidly passes off—*L* '97, i 1466, *B M J* '98, ii 619, *B M J E* '97, i 55, 75, 87, 92, '98, i 99, *Pr* lx 503

A 1 p c solution did not show the slightest cloudiness when allowed to stand in an open vessel for two months—*P J* '97, i 368

It is stated to possess the following advantages (1) it does not cause mydriasis, (2) does not affect accommodation, (3) causes deeper anaesthesia of the iris, (4) often proves efficient in cases of painful inflammation where Cocaine fails, (5) produces no toxic effects unless injected subcutaneously or swallowed, (6) has no effect on the corneal epithelium, (7) is strongly bactericidal in action. The solutions should be preserved in porcelain and not in glass vessels—*T G* '99, 322, 612, *B M J E* '99, ii 20, *Pr* lxiv 476, *M A* '00, 28

As a local anaesthetic in ophthalmic practice should not be used in stronger doses than 1 p c, as it is poisonous—*L* '05, ii 885

It is stated (*L* '06, ii 15) to be a most valuable addition to Cocaine in all operations in which it is necessary to cut the iris. The favourite combination recommended is Cocaine Hydrochloride, 2 p c, Holocaine Hydrochloride, 1 p c, dissolved in solution of Adrenalin Hydrochloride, 1 in 1000, and freshly prepared immediately before being used

ACOINE (Di-para anisyl-mono phenetyl guanidine hydrochloride)—A white, crystalline powder, soluble 1 in 50 of Water. Introduced as a substitute for Cocaine as being less toxic

A useful solution for producing anaesthesia is Acoine, 1, Sodium Chloride, 8, Distilled Water, 1000. Concentrated solutions should not be employed, as they give rise to irritation. The solutions, moreover, should not be exposed to the light—*L* '99, i 1372, *B M J* '99, i 1340, *P J* '99, i 538, *C D* '99, i 701

In subconjunctival injections as a local anaesthetic, no pain was produced when a mixture of equal parts of a 1 in 1000 solution of Mercury Cyanide and a 1 in 100 Acoine solution was injected—*L* '99, ii 1082

Solutions of 1 in 100 and 1 in 300 produce satisfactory anaesthesia in an unirritated eye, when there was much congestion it failed—*T G* '99, 697, *B M J E* '99, ii 76

NIRVANIN (Hydrochloride of Diethyl-glycocoll para-amido ortho-hydroxy-benzoic Methyl Ester)—Small white prisms, readily soluble in Water. Introduced as a local anaesthetic in surgical and dental operations. As a substitute for Cocaine and Orthoform, generally used in the form of a 2 p c solution. A 5 p c solution causes irritation when dropped into the eye. As much as 7 grains may be injected hypodermically without injury. A 1 p c solution has a marked bactericidal action—*P J* '99, i 95, 481, *C D* '99, i 701

NERVOCIDINE—A yellow, hygroscopic, amorphous powder, readily soluble in Water, slightly soluble in Alcohol (90 p c) and in Ether. It is obtained from an Indian plant 'Gasu Basu'. Introduced as a local anaesthetic. The irritation which it produces, the length of time required to produce anaesthesia, and its liability to produce toxic symptoms, however, preclude its general use. At present its employment is restricted to dental work—*L* '02, i 127, *P J* '02, ii 211

ALYPIN Primary Benzoyl-tetramethyldiamino- β -ethyl dimethylcarbinol Monohydrochloride $C_{18}H_{28}N_2O_2 \cdot HCl$, eq 312.39

A non-hygroscopic odourless crystalline powder

Solubility—1 in 1 of Water, 1 in $1\frac{1}{2}$ of Alcohol (90 p c), insoluble in Ether

Dose— $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.0064 to 0.032 gram

It was introduced as a local anaesthetic. It is stated (*L* '05, ii 321) to be easily absorbed by the mucous membranes and the subcutaneous tissue. It has a similar anaesthetic action to Cocaine. Solutions should always be freshly prepared, they may be sterilised by boiling without impairing their anaesthetic action

Enthusiastic account of it as a local anæsthetic. A 1 p c solution was used, it can be sterilised by boiling without spoiling, and it produces no bad effects, either general or local. Severe operations of considerable duration were performed with complete anæsthesia. The anæsthesia was given locally and never by lumbar puncture.—*B M J E* '07, ii 84

Nearly the equal of Cocaine in anæsthetic action, complete anæsthesia of the eye can be produced by a 1 or 2 p c solution in a minute.—*B M J E* '07, i 52

Tests—Alypin loses a small quantity of moisture when dried at a temperature of 100° C (212° F), the loss being equivalent to about 4.5 p c. The dried salt melts at about 170° C (338° F). It dissolves readily in Water, yielding a solution which is neutral in reaction towards Litmus paper. The aqueous solution yields with the usual alkaloidal precipitants, e.g., Potassiummercuric Solution, Iodo-potassium Iodide (Wagner's) Solution, Picric Acid Solution, etc. It also yields precipitates with Potassium or Sodium Hydroxide Solution and with Ammonia Solution, but is not precipitated by dilute Sodium Bicarbonate Solution. The solution acidified with diluted Nitric Acid yields on the addition of a solution of a white curdy precipitate insoluble in Nitric Acid, and if the precipitate be separated and washed, it is soluble in Ammonia Solution and Potassium Cyanide Solution. If a minute quantity of the salt be placed on the tongue it produces a characteristic sense of numbness resembling that produced by Cocaine. When ignited with free access of air the salt should leave no weighable residue.

NOVOCAINE—Para-aminobenzoyl-diethylamino ethanol Hydrochloride $C_{13}H_{20}N_2O_2.HCl$, eq 270.66

A white odourless crystalline powder which is readily soluble in cold Water. The aqueous solutions are neutral in reaction towards Litmus paper, and can be boiled without decomposition. It is introduced as a local anæsthetic, and is stated to be free from irritant action on living tissues. It is used chiefly in the form of solution, or as tablets, the powder having been withdrawn from the market. For hypodermic use the solutions vary in strength from a $\frac{1}{2}$ to 2 p c. For infiltration anæsthesia and for anæsthesia of thick layers of tissue, 1 to 2 p c isotonic solutions are used, for anæsthesia of the nerve-centres, 1 to 2 p c trunks 1 to 2 p c isotonic solutions are employed, for medullary anæsthesia a 5 p c isotonic solution has proved most useful. In ophthalmic practice 5 and 10 p c solutions are chiefly employed and produce no dilatation of the pupil or irritation, solutions of similar strength are employed in operations or explorations of the throat and nose, 10 to 20 p c solutions for anæsthesia of the larynx and pharynx, and in dental practice 1 to 2 p c isotonic solutions.

The dose for internal administration is $\frac{1}{2}$ to 1 gram.

Spinal and local anæsthetic. Advantages over Stovaine.—*B M J* '07, ii 876

85 cases, results uniformly good, should always be combined with Adrenalin.—*L* '07, ii 1686

160 cases, writer very well satisfied.—*F T* '07, 90

A satisfactory local and spinal anæsthetic.—*B M J E* '08, i 23

The safest and most perfect of known local anæsthetics in teeth extraction.—*B M J* '07, i 1172

STOVAINE—The Hydrochloride of α -methyl-2-methyl-prop-1-anol Benzoylamine $C_{14}H_{19}NO_2.HCl$, eq 269.63

Colourless, glistening, lamellar crystals, or a white, inodorous, crystalline powder.

Solubility.—1 in $1\frac{1}{2}$ of Water, 1 in 4 of Alcohol (90 p c), insoluble in Ether.

Dose.— $\frac{1}{2}$ to $\frac{1}{2}$ grain = 0.01 to 0.3 grammes

It is a spinal anæsthetic and said to be only half as toxic as Cocaine.

20 grains have been frequently used in the form of a 2 p c solution in distilled Water or physiological solution is the most useful, if a minimum of a 1 in 1000 Adrenalin solution be added to every 3 c.c., a still better result is obtained. Maximum dose of Stovaine is 40 to 50 grains, and of Adrenalin 0.001 gramme.—*B M J E* '05, i 92.

In work upon the nose, throat and ear, a 5 p c solution was found (*B M J E* '05, 11 43) to be equal in its effects to a 10 p c solution of Cocaine. As a local anæsthetic for the larynx Stovaine gave fair results. No appearances of poisoning have been observed after its application. $\frac{1}{2}$ to 1 p c aqueous solutions have been used by infiltration for the production of local anæsthesia. $\frac{1}{2}$ to $\frac{3}{4}$ p c solutions of Stovaine act as powerfully as 1 p c solutions of Cocaine—*B M J E* '05, 11 95.

The chief physiological difference from Cocaine is that it is a vaso dilator, not a vaso constrictor, and further, it seems to have a tonic effect upon the heart. Hence the vascular system seems to escape all the harmful effects of Cocaine. Of great value for the production of anæsthesia by intraspinal injection in acute general peritonitis. The minimum dose that can be given without fear of respiratory paralysis is 0.5 c c of a 10 p c solution. If the anæsthesia is required low down in the leg, quite a small dose, often as little as 0.3 c c of a 10 p c solution, is sufficient. In acute cases the method of gradual injection is imperative. A commencing dose of 0.6 c c may be given as a minimum, and if at the end of 7 minutes it is obvious that the dose is too small, an extra 0.3 c c should be given until the extent of the anæsthesia required is obtained—*B M J* '06, 1 1089.

Notes on 100 cases, an excellent anæsthetic though a little uncertain in its action—*B M J* '07, 11 14.

In 78 p c of the cases injected nephritis followed, and lasted from 6 $\frac{1}{2}$ to 30 days. The kidneys were previously healthy. Kidney mischief is therefore an absolute contra-indication—*B M J* '07, 11 1003.

For heavy men a dose of 7 to 8 cc is necessary, 5 to 6 for a person of medium weight. In most cases anæsthesia is obtained after five minutes, and lasts from three quarters to one hour. For operations on the abdominal wall it is advisable to make the puncture in the third lumbar space—*B M J E* '07, 11 56.

The minimal dose that can be given without fear of respiratory paralysis is 0.05 gramme. As quite a small amount of Sodium Carbonate renders Stovaine inactive, the Water in which the syringe is boiled ought not to contain any. Has a distinct influence in protecting from surgical shock. Method of administering, dosage, etc., fully considered—*B M J* '07, 11 869.

Barker finds a 5 p c solution in Distilled Water freezes at about -0.58°C , Blood Serum freezing at -0.56°C . If this were the only test, the solution should be isosmotic with the blood. If, however, a drop of blood is added to a little of a 4 or 5 p c solution of Stovaine, in five minutes the red blood corpuscles swell and become pale, and in ten minutes are almost invisible. In a really isotonic fluid, such as normal Saline (0.91 p c Sodium Chloride) or normal Glucose Solution (5 p c Glucose) the salts are seen unchanged in 24 or 48 hours. The formula suggested by Barker (*B M J* '07, 1 670) is Stovaine, 10 grammes, Glucose, 5 grammes, Sterilised Distilled Water, to produce 100 c c.

A second series of 100 cases of spinal analgesia in which this drug was employed is recorded by Barker—*B M J* '08, 1 248. The solution used contained 5 p c of Stovaine, and 5 p c of Glucose, in Water. It had a specific gravity of 1.023 as against 1.007 of the cerebrospinal fluid. The average amount of the solution usually injected was 1 c c = 0.05 gramme of Stovaine.

A report of 50 cases of analgesia produced by the intraspinal injection of Stovaine—*L* '08, 1 1058.

Suspicion expressed that some of the ill effects reported abroad after spinal analgesia were due to the Adrenalin principle added, and not to the anæsthetic drug at all—*B M J* '07, 1 665, '08, 244.

Disadvantages: the analgesia is not reliable and of short duration, the bowels are frequently opened on the operating table, may cause respiratory paralysis, many deaths and several cases of permanent paralysis have followed the spinal injection of Stovaine—*B M J* '07, 11 876.

Favourable results in 100 cases—*B M J* '07, 11 12.

Fatal paralysis in a man after injection of 0.05 gramme—*L* '07, 1 45.

Tests—Stovaine melts at 175°C (347°F). It dissolves readily in Water, forming a solution which is neutral in reaction towards Litmus paper. A weighed quantity of 0.5 gramme of Stovaine, evaporated with 1 c c of a mixture of equal parts of Hydrochloric and Nitric Acid on a water-bath, yields a colourless residue which has a pungent odour, and if to the residue 1 c c of Potassium Hydroxide

Solution be added and the mixture evaporated, the residue has a fruity odour and only drops separate on the addition of Water. Its aqueous solution is precipitated by the chief alkalis, Potassio-mercuric Iodo-potassium solution, Picric Acid, yields precipitates on the addition of Potassium or Sodium Hydroxide Solution, and on the addition of Ammonia Solution. The aqueous solution, when acidified with Nitric Acid, yields on the addition of Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, and which when separated and washed dissolves readily and completely in Ammonia Solution or in Potassium Cyanide Solution. 0.5 gramme of the salt when ignited with free access of air should leave no weighable residue.

COCCUS.

COCHINEAL

FR, COCHENILLE, GER, COCHENILLE, ITAL, COCCINIGLIA, SPAN, COCHINILLA

The dried fecundated female insect, *Coccus Cacti*, reared on *Nopalea cochenillifera*, and on other species of *Nopalea*

When dried in the sun the insects are of an ash-grey colour with a silvery surface, but when killed by immersion in boiling Water they have a reddish appearance, and if dried by artificial heat they are black

Used as a colouring agent

Official Preparation.—Tinctura Cocci. Used in the preparation of Tinctura Cardamomi Composita and Tinctura Cinchonæ Composita

Not Official.—Carmine, Glycerinum Carmini, Liquor Carmini, Liquor Coccineus, Glycerinum Cocci, Liquor Cocci, and Syrupus Coccionellæ

Foreign Pharmacopœias.—Official in US (*Coccus*), Jap and Swiss (*Coccionella*), FR (*Cochenille*), Port (*Cochonilha*), Mex and Span (*Cochinilla*). Not in the others

Descriptive Notes.—The dried insect forming this drug is named *Coccus Cacti*, Linn, in the *BP*, and *Pseudococcus Cacti*, Burmeister, in the *USP*. The dried insect is imported chiefly from Teneriffe, and is met with in commerce in four principal forms, known as silver grain, black grain, madres or zacatille, and granilla or siftings. The black grain consists of the insect dried by artificial heat, the silver grain is dried in the sun. The 'madres' consist of the female insect collected in March after the young are hatched, and the granilla consists chiefly of young insects sifted out. For the manufacture of Carmine the madres are preferred as, weight for weight, they yield more colour than before the young have left the mother. They are black and very concave on the under surface, not flat or slightly convex as in the ordinary silver and black grain. The insect is sometimes adulterated with Sulphate of Barium or of Lead and other mineral matter to increase its weight or improve its colour. It is sold at a lower price than the black variety which is more rarely adulterated, and then only with Iron Sand, which is visible under a good lens. The size is given both in *BP* and *USP* as 5 mm (1/4 in) long. *BP* states that it is somewhat oval in outline, flat or concave beneath, convex above, transversely wrinkled, purplish-black or purplish-grey, easily reduced to powder, which is dark-red or purple-coloured. *USP* describes it as

somewhat oblong and angular in outline, flat and concave beneath, convex above, externally purplish-grey or purplish-black, transversely wrinkled, easily pulverisable, yielding a dark-red powder

Tests—The *BP* requires that no insoluble powder should separate when the specimen is macerated in Water, and that the ash should not amount to more than 6 p c. The *USP* limit of ash is also 6 p c. The ash of Cochineal varies very much. Out of 44 samples examined in the author's laboratory 17 came within the *BP* limits, 3 yielded between 6 and 8 p c of ash, 23 exceeded 8 p c, and 1 sample contained as much as 37.4 p c of ash.

A comparison of the colouring power of the Cochineal may be made by powdering the sample with some broken glass, macerating for 24 hours in Water, with intervals of frequent stirring, and filtering through paper. The clear filtered liquid may be compared with a similar product prepared from a standard specimen.

Preparation

TINCTURA COCCI TINCTURE OF COCHINEAL

1 of Cochineal in powder, macerated with 10 of Alcohol (45 p c)
(1 in 10)

Dose—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Dan, 1 in 5, Fr and Mex, 1 in 10. By weight. Not in the others.

Tests—Tincture of Cochineal has a specific gravity of about 0.950, contains about 2.5 p c w/v of total solids and about 45 p c w/v of Absolute Alcohol.

Not Official

CARMINE—Prepared from Cochineal, an excellent colouring agent for powders and ointments. It is also used as a staining agent in microscopy.

GLYCERINUM CARMINI—Carmine, 3, Distilled Water, 3, Solution of Ammonia, *BP*, 4, dissolve and add gradually Glycerin, 18, heat in a water-bath till free from ammoniacal odour. When cold add solution of Ammonia, 1, to prevent gelatinisation, and Distilled Water, q s to 24.—*Martindale*

BPC is almost identical with this.

LIQUOR CARMINI—Carmine, 6, Water of Ammonia (10 p c), 35, Glycerin, 35, Water, q s to yield 100.—*USNF* 1896

This has been incorporated in the *BPC*, but in the *BPC Supplement* it may be made in a similar way to *Liquor Cocci*, replacing the Cochineal by 6 of Carmine.

USNF 1906 has altered the figures to 6.5, 36.5, 36.5 in 100.

LIQUOR COCCINEUS—Cochineal, in No. 50 powder, 6, Potassium Carbonate, 3, Alum 3, Potassium Bitartrate, 6, Glycerin, 50, Alcohol (90 p c), 3, Distilled Water, q s to yield 100.—*USNF* 1896

GLYCERINUM COCCI—Cochineal, unbruised, 20, Potassium Carbonate, 1, Potassium Citrate, 10, Glycerin, 20, Distilled Water, sufficient to produce 100.—*BPC*

LIQUOR COCCI—Cochineal, 20, Potassium Carbonate, 1, Potassium Citrate, 10, Alcohol, 20, Distilled Water, to produce 100. Dissolve the Potassium Carbonate in 60 of the Distilled Water and digest the unbruised Cochineal in the solution on a water bath for 6 hours or until exhausted, then strain, cool, add

the Alcohol and Potassium Citrate, and make up the required volume with Distilled Water — *B P C*

This is not an economical preparation, half the quantity of Cochineal will produce a solution of practically the same depth of colour. It is also an advantage to just bruise the Cochineal to a very coarse powder, but when not bruised it is necessary to assist the straining by gentle pressure, otherwise a considerable proportion of the liquid will be left in the marc. The process may be modified with advantage as follows — Cochineal (in coarse powder), 10, Potassium Carbonate, 1, Potassium Citrate, 10, Alcohol, 20, Distilled Water, *q s* to make 100. Dissolve the Potassium Carbonate in 60 of Distilled Water, and digest the Cochineal in the solution on a water-bath for 3 hours, replace the Water lost by evaporation, then strain and cool, add the Alcohol and Potassium Citrate, pass more Distilled Water through the strainer to make the total volume 100. Filter. This modification applies also to the Glycerinum Cocci, adding the Glycerin and Potassium Citrate to the strained fluid.

SYRUPUS COCCIONELLÆ — Cochineal, 10, Potassium Carbonate, 1, Rose Water, 150, Cinnamon Water, 150, digest for 4 hours, then filter, in each 100 parts of the filtrate dissolve Sugar, 160, and Alum, 0.1, boil and strain, all by weight — *Austrian Elenchus*

CODEINA.

CODEINE



A crystalline alkaloid (Methyl Morphine) obtained from Opium or synthetically from Morphine.

Solubility — 1 in 80 of Water, 1 in 24 of boiling Water, 1 in 2 of Alcohol (90 p c), 1 in 2 of Chloroform, 1 in 30 of Ether, 1 in 12 of Benzol, 1 in 85 of Liquor Ammoniac, 1 in 58 of Ether, sp gr. 0.720.

Medicinal Properties — It stops or lessens the glycosuria in diabetes (an entire abstinence from starchy food being strictly observed) in doses of 1 grain 3 times a day, gradually raised to 2 grains. Useful in relieving the hacking cough of phthisis, in ovarian pain, and as a mild hypnotic.

It has a powerful action in allaying abdominal pain, and it can be pushed to a much greater extent than Morphine without causing drowsiness or interfering with the respiration or with the action of the bowels — *B M J* '88, 1. 1214.

In $\frac{1}{4}$ to $\frac{1}{2}$ grain doses combined with Phenacetin has been stated to give relief from the severe headache and general paroxysms in malaria — *Pr* lxxii 682.

Dose — $\frac{1}{4}$ to 2 grains = 0.016 to 0.13 gramme.

Sus., maximum single dose, 0.1 gramme, maximum daily dose, 0.4 gramme.

Prescribing Notes — For coughs it is usually given in the form of lozenges or pastils, or as a pill, using Powder of Gum Acacia and 'Tr. U. S. G. A. S.' as excipients. For diabetes it is sometimes combined with Extract of Cascara. Codeine Phosphate in solution is used for hypodermic injection.

Official Preparations — Symplic Codeinæ from Codex Pharmacopœia.

Not Official — Codemæ Pastillæ, Tinctura Codemæ, Pilula Codeinæ Composita, Codeinæ Iodas, Apocodemæ Infusio.

Foreign Pharmacopœias — Official in Dan., Dutch, Fr., Hung., Ital., Mex., Port., Russ., Span., Swed., Swiss and U. S. Not in the others.

Tests—Codeine possesses a melting point of 156°C (312°F), the *BP* does not give a melting point, the *USP* gives 154°C (310°F), *Fr. Codex* (1908) 155°C (311°F). The aqueous solution of the alkaloid is alkaline towards Litmus paper and is laevogyrate. Pure Sulphuric Acid yields no coloration with Codeine in the cold, but on warming a blue coloration is slowly produced, the blue coloration is immediately produced if a trace of Ferric Chloride, Ammonium Molybdate Solution, or Potassium Ferricyanide is present. The *BP* adds that on the addition of a minute trace of Diluted Nitric Acid the colour changes to a bright scarlet, becoming orange. Codeine is distinguished from Morphine by the following tests—When moistened with Nitric Acid the liquid becomes yellow, but not red. A 1 in 50 solution of Codeine in Water, acidulated with Hydrochloric Acid, yields with Potassium Hydroxide Solution a whitish precipitate, but is not precipitated by the addition of Ammonia Solution. A saturated aqueous solution acidulated with Hydrochloric Acid should yield on the addition of Ferric Chloride Test-solution and a very dilute Potassium Ferricyanide solution only gradually a dirty green, but no blue coloration. The *BP* wording of the latter test has been subjected to criticism (*PJ* '00, ii 149). As the text reads a neutral solution of Codeine Hydrochloride would be used in making this test, which would obviously be quite different in its character from the solution employed if a comma were placed after the word Water, so as to read 'A saturated solution of Codeine in Water, acidulated with Hydrochloric Acid'. Exception is taken in the same reference to the *BP* statement that the test shows the absence of Morphine and other impurities. This is considered too broad as it will by no means detect all other impurities nor even any considerable proportion of possible impurities.

0.1 gramme when heated to redness with free access of air leaves no weighable residue.

Codeine is not official in the *PG*.

Colour Reactions—The following colour reactions are given in the *BP* and *USP*. Sulphuric Acid dissolves the alkaloid without coloration, *BP*, either without coloration or producing a slight pinkish tint which disappears within two minutes, but on heating a violet colour is developed. (The presence of nitrous compounds causes a pink colour in the cold), *USP*. The solution of Codeine in Sulphuric Acid gives a blue or bluish-black colour with (a) 2 drops of T.S. of Ammonium Molybdate, (b) a trace of Ferric Chloride, (c) or Potassium Ferricyanide, the addition of a minute trace of Diluted Nitric Acid changes this colour to a bright scarlet, becoming orange, *BP*. Codeine yields with Sulphuric Acid and (a) a trace of Ferric Chloride, a violet-blue coloration, (b) a drop of Nitric Acid, a blood red coloration, (c) a trace of Selenious Acid, a green coloration, (d) a drop of T.S. of Formaldehyde (added to the Codeine and Sulphuric Acid previously mixed), a violet-blue coloration, *USP*. A mixture of Codeine and Nitric Acid yields a yellow coloration, but should not be red, *BP*, the *USP* states that if 0.05 gramme of Codeine be sprinkled upon 2 c.c. of Nitric Acid (sp. gr. 1.200) the crystals will turn red, but the acid will only acquire a yellow colour (difference from and absence of Morphine). A solution of 0.08 gramme of Codeine in 2 c.c. of Sulphuric Acid yields with 1 drop of a diluted Nitric Acid Solution (1 drop of Acid in 200 c.c. of Water), a bluish-red tint, gradually changing to blue, *USP*. There should not be any blue colour developed, but only slowly a dull green on the addition of T.S. of Ferric Chloride.

and a very dilute solution of Potassium Ferricyanide to a saturated solution of Codeine in Water acidulated with Hydrochloric Acid, *BP*. In the *USP* test quantities are given a small crystal of Potassium Ferricyanide is dissolved in 10 c.c. of Water and to this is added 1 drop of T.S. of Ferric Chloride and 1 c.c. or a 1 p.c. solution of Codeine, when no blue colour should be developed at once.

CODEINÆ PHOSPHAS. CODEINE PHOSPHATE

This crystalline product ($C_{18}H_{21}NO_3, H_3PO_4)_2, 3H_2O$, eq 842 20, is the most soluble salt of Codeine.

Fine white acicular crystals, or as a white odourless crystalline powder possessing a bitter taste and feebly acid reaction.

Solubility—1 in 4 of Water, 1 in 200 of Alcohol (90 p.c.)

Dose.— $\frac{1}{4}$ to 2 grains = 0.016 to 0.13 gramme

Ph. Ger. maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme

Foreign Pharmacopœias—Official in Belg., Dan., Fr., Ger., Jap., Norw., Russ., Swed., Swiss and U.S.

Tests.—Codeine Phosphate should answer the tests distinctive of Codeine given in the large type under the heading of Codeina. It melts at a temperature of about $235^{\circ}C$ ($455^{\circ}F$). The aqueous solution yields on the addition of Silver Ammonio-nitrate Solution a light yellow precipitate readily soluble in Ammonia Solution and in cold dilute Nitric Acid, with Magnesium Ammonio-sulphate Solution it affords a white crystalline precipitate, with Ammonium Methyl containing Nitric Acid, it yields on warming a yellow precipitate, which is soluble in Ammonia and which is reprecipitated as a white crystalline precipitate on the addition of Magnesium Ammonio-sulphate. The reaction of a 5 p.c. aqueous solution towards Litmus paper is slightly acid. This solution yields with Potassium Hydroxide Solution a whitish precipitate, but no precipitate on the addition of Ammonia Solution.

Codeine Phosphate of the *BP* contains theoretically 70.52 p.c. of Codeine, *USP* 69.05 p.c. The *BP* gives no process for quantitatively determining the amount of alkaloid present, the *USP* requires that a weighed quantity of 0.2 gramme of the salt when precipitated with Potassium Hydroxide Solution and shaken out with Chloroform should yield not less than 0.13 gramme of Codeine, corresponding to not less than 65.0 p.c. of alkaloid.

The more generally occurring impurities are excess of moisture, Chlorides, Sulphates, Morphine and mineral matter. The formula given for the salt in the *BP* shows $1\frac{1}{2}H_2O$, whilst that of the *USP* and the *Fr. Codex* (1908), indicates $2H_2O$. All three Ph. state that at $100^{\circ}C$ ($212^{\circ}F$) it loses all of its Water of crystallisation. The *BP* formula $1\frac{1}{2}H_2O$ would indicate a loss of 6.36 p.c. of Water, the *USP* and *Fr. Codex* a loss of 8.31 p.c., the *PG* mentions that at $100^{\circ}C$ ($212^{\circ}F$) the salt loses 8 p.c. in weight. The aqueous solution when acidified with diluted Nitric Acid should yield no opalescence on precipitation with Silver Nitrate Solution or with Barium Chloride Solution indicating the absence of Chlorides and Sulphates. The presence of Morphine is indicated by Ferric

Chloride Test-solution which would yield a blue coloration The *USP* and *Fr Codex* employ a mixture of Potassium Ferricyanide Solution and Ferric Chloride Test solution as a test for Morphine

0.1 gramme of the salt leaves no weighable residue when ignited with free access of air

Colour Reactions—0.01 gramme of Codeine Phosphate gives with 10 c.c. of Sulphuric Acid a colourless solution, *BP*, the *USP* states that the acid produces either no colour or a slight pinkish tint which disappears in 2 minutes Codeine Phosphate gives with Sulphuric Acid containing (a) a trace of Ferric Chloride (1 drop of T.S. of Ferric Chloride in 10 c.c., *PG*) a violet-blue colour, *PG* and *USP*, (b) a trace of Selenious Acid, a green colour changing rapidly to blue and then slowly back to grass green (Morphine gives a blue colour changing to green and then to brown), *USP*, (c) a drop of T.S. of Formaldehyde, a violet blue colour (Morphine gives an intense purple), *USP* The solution of a small crystal of Potassium Ferricyanide in 10 c.c. of Water with the addition of a drop of T.S. of Ferric Chloride should not immediately assume a blue colour when mixed with 1 c.c. of a 1 p.c. Solution of Codeine Phosphate, *PG* and *USP* No blue colour should be developed with Codeine Phosphate and T.S. of Ferric Chloride, *BP*

Codeinæ Hydrochloridum is official in Austr., Dutch and Mex. It is crystalline and soluble in Water. **Codeinæ Salicylas** is also crystalline, and readily soluble in Alcohol and Ether, but only slightly soluble in Water. **Codeinæ Sulphas** is official in U.S. Doses same as Phosphate

Preparation

SYRUPUS CODEINÆ SYRUP OF CODEINE

Codeine Phosphate, 40 grains, Distilled Water, $\frac{1}{4}$ fl. oz., Syrup, $19\frac{1}{2}$ fl. oz. (1 grain in 240 minims)

BP directs the Codeine Phosphate to be dissolved in the Distilled Water, but 40 grains of Codeine Phosphate will not dissolve in $\frac{1}{4}$ fl. oz. Distilled Water; it is better to use 180 minims

Dose— $\frac{1}{2}$ to 2 fl. drms = 1.8 to 7.1 c.c., containing $\frac{1}{8}$ to $\frac{1}{2}$ grain Codeine Phosphate

It is 50 p.c. stronger than the Syrup described in previous editions of the *Companion*

Foreign Pharmacopœias—Official in Belg., 9 in 1000, *F1*, Ital., Mex. and Swiss, 1 in 500, Span., 1 in 600. Made with the Alkaloid. Dutch, 1 of Hydrochloride in 400

Not Official

CODEINE PASTILS—Contain $\frac{1}{8}$ grain = 0.008 gramme of Codeine in each. One for a dose when the cough is troublesome. An improvement on Codeine Jelly

Official in Ital., $\frac{1}{8}$ grain = 0.005 gramme in each

LINCTUS CODEINÆ—Syrup of Codeine, 20 minims, Glycerin, 20 minims, Lemon Juice, 18 minims, Chloric Ether, 2 minims—*Brompton*
Syrup of Codeine, 80 minims, Citric Acid, 1 grain, Emulsion of Chloroform, 8 minims, Glycerin, 10 minims, Mucilage of Tragacanth, to 1 fl. drm—*St. Thomas's*

This has been incorporated in the *BP C*

Syrup of Codeine, $\frac{1}{2}$ fl. drms, Syrup of Virginian Prune, $\frac{1}{2}$ fl. drms—*Guy's*

PILULA CODEINÆ COMPOSITA—Codeine, $\frac{1}{2}$ grain, Kaolin, $\frac{1}{2}$ grain, Extract of Cascara, 2 grains, Hard Soap, to 4 grains—*Guy's*

CODEINÆ IODAS—A combination of Iodic Acid with the alkaloid. Has been introduced as an analgesic.

Dose.— $\frac{1}{2}$ grain = 0.082 gramme by hypodermic injection.

Apocodema—Produced by heating Codeine with Zinc Chloride, it forms brown amorphous resinous masses

Apocodemæ Hydrochloridum is supplied as a brown, amorphous powder, soluble in Water. Dott doubts the existence of Apocodeme, and states that the commercial products sold under this name are not of a very definite nature

It has been used by subcutaneous injection in 30-minim doses of a 1 p c solution to produce increased peristalsis of the bowel, and has also been used internally as an expectorant in bronchial affections and as a sedative in mental disturbance, in doses of 0.02 to 0.06 gramme ($\frac{1}{2}$ to 1 grain)

A suitable combination for the internal administration is Apocodeme Hydrochloride, 0.5 gramme, Syrup of Raspberry, 25 grammes, Distilled Water, 100 grammes, in doses of $\frac{1}{2}$ to 1 fl dr = 1.8 to 3.6 c c

This salt, which has been frequently referred to in medical literature, is again mentioned (L '06, 1 1191) as a laxative. It may be administered in doses of 2 c c of a 3 p c solution

COLCHICUM.

FR, COLCHIQUE, GER., ZEITLOSENKNOLLEN, ITAL, BULBO DE COLCHICO, SPAN, BULBO DE COLQUICO

The fresh Corm of *Colchicum autumnale*, as well as the dried, ripe Seeds, are official

The use of Colchicum Seeds only, was agreed by the Brussels International Congress

Medicinal Properties—It is a specific in gout, and will cure the acute form, controlling the pain and inflammation, and preventing the attack. Useful in the headache, dyspepsia, eczema, and other conditions which occur in gouty subjects. May be combined with other or given with saline purgatives in cases of hepatic congestion in gouty patients. It may produce gastric or intestinal irritation, even in ordinary doses, and should then be discontinued for a time. The Extract is frequently prescribed with Dover's Powder to relieve painful gout

Dose—Of the dried Corm, 2 to 5 grains = 0.13 to 0.32 gramme, but usually given in the form of Extract or Wine

Incompatibles—Tincture of Iodine, Guaiacum, and vegetable astringents

Official Preparations—Extractum Colchici and Vinum Colchici from the Corm, Tinctura Colchici Seminum from the Seeds

Not Official—Extractum Colchici Aceticum, Extractum Colchici Cormi, Fluidextractum Colchici Seminum, Mistura Colchici, Pilule Colchici et Hydrargyri, Tinctura Colchici Composita, Tinctura Colchici Florum, Vinum Colchici Seminum, Colchicina, and Colchicinæ Salicylas

Antidotes—In case of poisoning with Colchicum, emetics and demulcent drinks, and, if coma be present, Brandy, Ammonia, and powerful stimulants may be given. Hypodermic injection of $\frac{1}{2}$ c c of a 3 p c solution

COLCHICI CORMUS. COLCHICUM CORM

The fresh Corm of *Colchicum autumnale*, collected in early summer, also the Corm, dried at a temperature not exceeding 65° C (150° F), after being stripped of its coats and transversely sliced.

Colchicum Corms contain about 0.5 p.c. of Colchicine, but the *B.P.* makes no requirement that they shall contain a definite percentage of alkaloid, the *U.S.P.* requires that they shall yield not less than 0.35 p.c. of Colchicine, and indicates a method of determination.

Foreign Pharmacopœias—Official in Mex., Port. and U.S. Not in the others.

Larger equivalent doses of the corm than of the wine or tincture can be given without undesirable effects, and the powdered corm is stated to give better results in acute gout.—*B.M.J.* '04, ii 1460

Descriptive Notes—It should be noted that the extract is prepared from the fresh corm collected in early summer. *Colchicum autumnale*, L., is a very local plant in this country, and there is therefore some difficulty in obtaining the fresh corm. The best period is in July when the leaves have turned yellow or in August when the plant is in flower, as the corm is then in mature condition. According to Schroff's experiments the corms are best dried entire in sun and air, and preserved, they lose none of their activity even if kept several years (*Pharmacographia*). Although the fresh corm is, according to the *B.P.* an inch broad (25 mm.) and an inch and a half long (35 mm.) it is often larger if obtained from full grown plants, and even the dried corm in slices may exceed an inch in diameter and $\frac{1}{10}$ to $\frac{1}{8}$ in thickness (2 or 3 mm.). The reniform shape is characteristic, for although the corms of some Fritillaries and the Hermodactyls of the East have the same shape they are not met with in Western commerce. Under the microscope the compound starch grains (0.1 to 0.15 mm., *Planchon* and *Collin*), usually three or four, each with a stellate hilum, and without stræ, and the irregular epidermal cells with pitted walls, are sufficiently characteristic. According to Vogl the cambium tissue contains a yellowish amorphous substance which, when a section of the corm is treated with concentrated Sulphuric Acid, colours the cellulose tissue gamboge yellow and the vessels orange red. The dried corm if long kept and especially if allowed to become damp loses its medicinal effect to a certain extent.

Tests—The *U.S.P.* process of determination is essentially as follows.—A weighed quantity of 10 grammes is macerated for twelve hours, with frequent intervals of shaking (or for four hours if a mechanical shaker be employed), in an Erlenmeyer flask with 100 c.c. of a mixture of 77 c.c. of Ether, 25 c.c. of Chloroform, 8 c.c. of Alcohol (94.9 p.c.), and 3 c.c. of Ammonia Water. The liquid is filtered, and 50 c.c. of the filtrate is evaporated to dryness at a gentle heat, the residue is dissolved in 10 c.c. of Ether, 5 c.c. of Water added, the mixture well stirred and the Ether evaporated. When cool the aqueous solution is filtered into a small separator, the insoluble matter being kept as much as possible in the beaker or dish. The residue is redissolved in Ether, 5 c.c. of Water added, the Ether evaporated as previously. The beaker or dish and the filter are washed with a little Water, and the combined aqueous solutions are well shaken with 15 c.c. of Chloroform. After separation of the chloroformic liquid, the aqueous portion is shaken with three

successive portions of 10 cc each of Chloroform, the chloroformic layer being separated in each case, mixed with the first Chloroform shaking, and the mixed chloroformic liquids evaporated to dryness. The residue is dissolved in a little Alcohol (94.9 pc), the Alcohol evaporated, the residue dissolved in 5 cc of Ether, 5 cc of Water added, and the mixture stirred for a few seconds. The Ether is evaporated, the aqueous liquid filtered through a wet filter paper into a separator, the vessel and filter washed with 5 cc of Water, the washings being added to the contents of the separator. The aqueous liquid is shaken out with 15 cc of Chloroform, the Chloroform transferred after complete separation to a tared flask, the aqueous portion extracted with three successive portions each of 10 cc of Chloroform, which are separated as previously and transferred to the tared flask containing the first shaking. The Chloroform is completely removed by evaporation, the residue is dissolved in Alcohol (94.9 pc), which is in turn evaporated, and the residue is dried at 100° C (212° F) until constant. If this weight be multiplied by 20 it indicates the pc of Colchicine present in the Corms.

Preparations

EXTRACTUM COLCHICI. EXTRACT OF COLCHICUM

A soft extract prepared from the juice of fresh *Colchicum Corms* which have been deprived of their coats. The clarified juice is heated to 100° C (212° F) to coagulate Albumen, and the strained liquid is evaporated to a soft extract at a temperature not exceeding 71.1° C (160° F).

100 lb of Corms yield about 4 lb of Extract.

The *BP* Extract is not a standardised preparation and no process of determination appears in the *Pharmacopœia*. The Extract official in the *USP* is standardised to contain 1.4 pc of Colchicine. The *PG* does not contain an Extract of Colchicum Corms.

Dose.— $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Foreign Pharmacopœias—Official in Belg, Fr, Ital, Mex and Span (Alcoholic Extract of Seeds), Mex (Alcoholic from Corms), also (Fluid Extract of Corms and Seeds), Port and US Extract from Corms with Acetic Acid. US has Fluid Extract of Seeds. Not in the others.

Tests.—The *USP* method for the determination of Colchicine in Colchicum Extract is essentially as follows.—A volume of 4 grammes of the extract is dissolved in 20 cc of Water, transferred to a 100 cc flask and sufficient Alcohol (94.9 pc) added to bring the volume of the liquid to 100 cc. After the contents of the flask have been well shaken, they are allowed to stand 5 minutes, filtered, and 50 cc of the filtrate collected, evaporated to dryness in a porcelain dish, 10 cc of Ether and 5 cc of Water are added to the residue, and the Ether removed by evaporation. When cool, the aqueous solution is filtered into a separator, the insoluble matter being retained as far as possible in the vessel used for the evaporation. The residue remaining in this vessel is again treated with 10 cc of Ether and 5 cc of Water, the Ether being removed as previously. The dish and filter are rinsed with a little Water, and the

mixed aqueous liquids collected in a separator After rendering the liquid alkaline by the addition of a sufficiency of Ammonia Solution, the Colchicine is extracted by successive quantities of 20, 15 and 10 c c of Chloroform respectively The separated chloroformic solutions are mixed and evaporated to dryness, and the residue is mixed with two successive small quantities of Alcohol (94.9 p c), the latter being removed in each case by evaporation 5 c c of Water and 10 c c of Ether are added to the residue, the liquid shaken and the Ether evaporated When cool, the aqueous liquid is filtered into a separator, the flask and filter being rinsed with Water The Colchicine is extracted by successive shakings with 20 c c, 15 c c and 10 c c of Chloroform The separated chloroformic liquids are mixed, transferred to a tared flask, evaporated to dryness, the residue dissolved in 2 small successive quantities of Alcohol (94.9 p c), the latter removed by evaporation and the residue dried till constant at 100° C (212° F) The weight multiplied by 50 shows the p c of Colchicine present in the Corm Extract

VINUM COLCHICI COLCHICUM WINE

4 of Colchicum Corm in No 20 powder, macerated in 20 of Sherry
(1 in 5)

Dose—10 to 30 minims = 0.6 to 1.8 c c

Diluted Acetic Acid appears to be about as good a solvent as Sherry, but Alcohol (45 p c) was better than either —*P J* '97, i 173 Further notes on the same —*P J* '98, i 131

Foreign Pharmacopœias—Official in Port, 1 and 10 Madeira, Mex, 1 in 10 Sherry See also Vinum Colchici Seminum

Tests—Colchicum Wine possesses a specific gravity of about 1.013, contains about 8.5 p c w/v of total solids and about 20 p c w/v of Absolute Alcohol

The Wine official in the *U S P* is prepared with a standardised fluid extract made from Colchicum Seeds, and should contain 0.04 p c w/v of Colchicine

COLCHICI SEMINA COLCHICUM SEEDS

FR, SEMENCE DE COLCHIQUE, GER, ZWITLOSENSAMEN, ITAL, SEMI DI COLCHICO, SPAN, SEMILLA DE COLQUICO

The dried ripe seeds of *Colchicum autumnale*, L

The Seeds are official in *B P*, *U S P* and *P G* They usually contain 0.6 to 1.0 p c of Colchicine, but neither the *B P* nor the *P G* requires them to yield a definite percentage of alkaloid nor includes a method of determination The *U S P* requires that they shall yield not less than 0.45 p c of Colchicine

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Span, Swed, Swiss and U S

The *Brussels Conference* recommends only the Colchicum seed

Descriptive Notes—Colchicum seeds are pale brown and inodorous when freshly gathered, but become darker in drying and exude a saccharine matter consisting of Glucose The percentage of this varies, and must be taken into consideration in estimating the extractive of galenical preparations made with them The seeds are

extremely hard and tough and difficult to powder. They have a bitter acid taste, and are nearly spherical, about $\frac{1}{16}$ to $\frac{1}{8}$ inch (2 to 3 mm) in diameter, but pointed by a strophiole appendage at the hilum. The diameter given in *BP* is 2.5 mm, *USP* 2 mm, *PG* 3 mm. The seeds ripen in June.

Tests—The method of determination recommended by the *USP* is essentially as follows.—A weighed quantity of 10 grammes is macerated, with frequent intervals of shaking, for 12 hours (or for 4 if a mechanical agitator is employed), in an Erlenmeyer flask, with 100 cc of a mixture of 77 cc of Ether, 25 cc of Chloroform and 8 cc of Alcohol (94.9 p.c.), and 3 cc of Ammonia Solution. A measured quantity of 50 cc of the filtered liquid is transferred to a beaker or dish and evaporated nearly to dryness. The residue is dissolved in 10 cc of Ether, 5 cc of Water added, the mixture stirred well, and the Ether evaporated. When cool the aqueous liquid is filtered into a separator, the insoluble matter being retained as largely as possible in the beaker. This residue is redissolved in Ether, and after the addition of 5 cc of Water, the previous operation is repeated. The beaker and filter are washed with a little Water and the Colchicine extracted from the mixed aqueous liquids by agitation with 15 cc of Chloroform. The agitation is thrice repeated with successive quantities each of 10 cc of Chloroform. The chloroformic liquids are separated, transferred to a tared flask, the Chloroform completely removed by evaporation, the residue twice dissolved in small successive quantities of Alcohol (94.9 p.c.), the latter being in each case removed by evaporation, and the residue finally dried at a temperature of 100° C (212° F) till constant in weight. This weight multiplied by 20 gives the percentage of Colchicine present in the seeds. The ash of Colchicum Seeds should not exceed 6.0 p.c.

The *USP* has a 1 in 1 Fluid Extract prepared with Colchicum Seeds and standardised to contain 0.4 p.c. w/v of Colchicine.

Preparation

TINCTURA COLCHICI SEMINUM. TINCTURE OF COLCHICUM SEEDS

1 of Colchicum Seeds, in No. 30 powder, percolated with Alcohol (45 p.c.), to yield 5. (1 in 5)

BP 1885 was 1 in 8, altered in *BP* 1898 to 1 in 5, see also Foreign Pharmacopœias given below.

Dose.—5 to 15 minims = 0.3 to 0.9 c.c.

Ph. Ger. maximum single dose, 2.0 grammes, maximum daily dose, 6.0 grammes.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Jap., Span. and Swiss 1 in 10, Hung., 1 in 5, Ger., Ital. and Swed., 1 and 10, U.S., 1 in 10, all from Seeds. Mex., 1 in 5 with Corms. Port., 1 in 5 with Seeds and Corms. All by weight except U.S. Not in the others.

The *Brussels Conference* recommended a strength of 10 p.c. of the Seeds with Alcohol (70 p.c.). Belg., Dan., Fr. and Swiss adopt the *BC* Standard.

Tests.—Tincture of Colchicum Seeds possesses a specific gravity of from 0.950 to 0.955, contains from 1 to 3 p.c. w/v of total

solids and about 43 p c w/v of Absolute Alcohol The Tincture official in the *BP* is not a standardised preparation, and consequently no process is given for the quantitative determination of the Colchicine

The *USP* Tincture is required to contain 0.04 p c w/v of Colchicine A measured quantity of 100 c c is evaporated on a water-bath to about a tenth of its volume Sufficient Alcohol (94.9 p c) is added to redissolve any separated matter, 1 c c of Ammonia Solution added, and the Colchicine removed by shaking three times in succession with separate portions of 15 c c, 15 c c and 10 c c of Chloroform The Chloroform solutions are separated, mixed and evaporated nearly to dryness The residue is dissolved in 10 c c of Ether, and 5 c c of Water added, the mixture well stirred, and the Ether removed by evaporation When cool the aqueous solution is filtered into a small separator, the insoluble matter being retained as far as possible in the beaker This residue is redissolved in a little Ether, 5 c c of Water added, the process repeated as previously The beaker and filter are washed with a little Water, and the alkaloid is extracted from the aqueous solution by agitation, first with 15 c c, and thrice subsequently with successive quantities of 10 c c each of Chloroform The chloroformic liquids are separated in each case, transferred to a tared flask, the Chloroform removed completely by evaporation, the residue twice dissolved in a little Alcohol (94.9 p c), the latter being in each instance removed by evaporation, and the residue ultimately dried at a temperature of 100° C (212° F) till constant in weight This weight will represent the percentage w/v of Colchicine present in the Tincture

Not Official

EXTRACTUM COLCHICI ACETICUM (*BP* 1885)—Crushed fresh Corms, previously peeled, 112, Acetic Acid, 6, stir together, press, and after subsidence heat the clear liquor to 212° F (100° C), strain through flannel, and evaporate at 160° F (71.1° C) to a soft extract

100 lb of Corms yield about 5 lb of Extract

Dose— $\frac{1}{2}$ to 2 grains, in pill, with an equal weight of Liquorice Powder

EXTRACTUM COLCHICI CORMI (*US*)—1000 grammes of Colchicum Corms in No. 60 powder is percolated with 350 c c of Acetic Acid (56 p c) mixed with 1500 c c of Water, and the percolation completed with Water, the liquor is evaporated at a temperature not exceeding 80° C (176° F) in a porcelain vessel to a pulular consistence It is adjusted to a strength of 1.4 p c of Colchicine by means of Sugar of Milk

FLUIDEXTRACTUM COLCHICI SEMINIS—About a 1 in 1 Fluid Extract prepared from Colchicum Seeds, and a mixture of Alcohol (95 p c) 2 parts and Water 1 part It is standardised to contain 0.4 p c of Colchicine.—*USP*

MISTURA COLCHICI—Colchicum Wine, 15 minims, Potassium Bicarbonate, 20 grains, Magnesium Sulphate, 15 grains, Peppermint Water, to 1 fl oz —*St Thomas's*

The *BPC* formula is the same, except in using Magnesium Carbonate, 10 grains, in place of Potassium Bicarbonate, 20 grains

Colchicum Wine, 15 minims, Carbonate of Magnesia, 10 grains, Bicarbonate of Potassium, 15 grains, Peppermint Water, to 1 fl oz —*Royal Free*

PILULÆ COLCHICI ET HYDRARGYRI—Acetic Extract of Colchicum, $\frac{1}{2}$ grain, Mercury Pill Mass, $\frac{1}{2}$ grain, Compound Extract of Colocynthis, $\frac{1}{2}$ grain, to make one pill —*BPC*

Sir Benjamin Brodie's Gout Pills—Compound Extract of Colocynth, 16 grains, Extract of Rhubarb, 16 grains, Mercury Pill, 16 grains, Acetic Extract of Colchicum, 6 grains, divided in 12 pills—*Pharm. Form*

This has been incorporated in the *B P C* under the title *Pilulæ Colchici et Hydrargyri Compositæ*

TINCTURA COLCHICI COMPOSITA (*Ph. Lond.*)—1 of bruised Colchicum Seeds, macerated with 8 of Aromatic Spirit of Ammonia

Dose—15 to 30 minims = 0.9 to 1.8 c c

This has been incorporated in the *B P C*

TINCTURA COLCHICI FLORUM (*Squæ*)—Fresh Flowers, 2, Alcohol (90 p c), by weight, 1, after seven days, filter

B P C has incorporated this preparation, but employs Alcohol (70 p c) by volume in place of Alcohol (90 p c) by weight

Dose—10 to 30 minims = 0.6 to 1.8 c c This preparation closely resembles the *Lau. Medicinalis*

Tests—Tincture of Colchicum Flowers possesses a specific gravity of about 0.970, contains about 4 p c w/v of total solids and about 39 p c w/v of Absolute Alcohol. It is standardised to contain 0.06 p c of Colchicine

VINUM COLCHICI SEMINUM—1 of Colchicum Seeds, in fine powder, macerated with 10 of Sherry

Dose—10 to 30 minims = 0.6 to 1.8 c c

Ph. Ger. maximum single dose, 2 grammes, maximum daily dose, 6 grammes
B P C same strength as above, but with Detannated Sherry

Foreign Pharmacopœias—Official in Dutch, 1 and 10 M^g and 5 and 10 M^g, Hung, 1 in 5 and Norw, 1 and 10 Sherry, Ital, 1 and 10 Marsala, Port, 1 and 10 Madeira, U.S., Fluid Extract, 1 in 10, with White Wine and Alcohol. All by weight except U.S.

COLCHICINA Colchicine $C_{22}H_{25}NO_6$, eq 396.24—A yellowish powder, soluble in Water and Alcohol (90 p c)

Dose— $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.00054 to 0.002 grammes

Colchicine is Tri-methyl-acetyl-colchicemic Acid, and crystallises in shining white needles. Colchicine is the Methyl Ester of Colchicemic

Colchicine has been shown (*B M J* '04, ii 1697, *L* '04, ii 1784) to be a slow poison, acting on the medulla, which it paralyzes

A solution of $\frac{1}{10}$ grain in Methyl Salicylate enclosed in Gelatin capsule is known under the name of 'Colchisal'

Foreign Pharmacopœias—Official in Fr, Hung, Mex and U.S.

Tests—Colchicine melts at 148° to 147° C (289.4° to 296.6° F) *Fr. Codex* gives 145° C (293° F). It is neutral in reaction towards Litmus. Its aqueous solution is laevogyrate. Its solution in dilute mineral acid gradually becomes meniscus yellow. Concentrated Nitric Acid produces, on the colour changing to yellow, and ultimately to green, on dilution with Water the violet solution changes to yellow, and on the addition of Sodium Hydroxide Solution in excess, to a fine orange or red colour, with very minute quantities of Colchicine the colour produced is rose-red. In concentrated Sulphuric Acid the alkaloid dissolves with the production of an intense yellow colour, which changes to a greenish-blue on the addition of Nitric Acid. It then turns to red, and finally to yellow. An aqueous solution gives no immediate coloration with Ferric Chloride. But on warming a garnet-red coloration is produced, an alcoholic solution gives a garnet-red coloration. Colchicine is stated to form a compound with Chloroform which is readily decomposed by Water. *Fr. Codex* gives the formula of this compound as $C_{22}H_{25}NO_6 \cdot 2CHCl_3$. The absence of Chloroform may be ensured by mixing 0.1 gramme of the alkaloid with 0.3 gramme of Calcium Chloride (free from Chlorides) moistening with Water, evaporating to dryness, and igniting. The residue is dissolved in dilute Nitric Acid filtered, and Silver Nitrate Solution added, no turbidity or precipitate should be produced.

0.1 gramme of Colchicine should leave no weighable residue when ignited with free access of air

COLCHICINÆ SALICYLAS—Yellow, amorphous powder, soluble in Water, in Alcohol (90 p c), and in Ether. It dissolves 1 in 100 of Methyl Salicylate. It should be preserved in well-closed bottles of a dark amber tint. Has been recommended in the treatment of gout and rheumatism as combining the properties of its two constituents.

Dose— $\frac{1}{10}$ to $\frac{1}{32}$ grain = 0.00054 to 0.002 gramme

Tests—Colchicine Salicylate yields the reactions distinctive of Colchicine given under that heading.

An aqueous solution of the salt yields a violet coloration with Ferric Chloride Test Solution. The salt should leave no weighable residue when ignited with free access of air.

Not Official

COLLINSONIA

The Root of *Collinsonia Canadensis*, L. (Stone Root)

Various preparations of this have been recommended in acute cystitis, and in the treatment of renal calculi.—*B M J* '87, 11 712, *L* '88, 1 868

Dose—15 to 60 grains = 1 to 4 grammes

TINCTURA COLLINSONIÆ—Collinsonia Root, 1, Alcohol (60 p c), 10

Dose—30 to 120 minims = 1.8 to 7.1 c c

This has been incorporated in the *B P C*

A 1 in 1 Fluid Extract is also made

Dose—15 to 60 minims = 0.9 to 3.6 c c

COLOCYNTHIDIS PULPA.

COLOCYNTH PULP

FR, COLOQUINTE, GER, KOLOQUINTHEN, ITAL, COLOQUINTIDE,
SPAN, COLOQUINTIDA

The dried pulp of the Fruit of *Citrullus Colocynthis*, Schrader, freed from Seeds. The fruit is imported chiefly from Smyrna, Trieste, France and Spain.

Medicinal Properties—It is a powerful drastic, hydiagogue cathartic, dangerous in large doses. It should not be prescribed alone, but in combination it is very commonly prescribed as an aperient in the form of Compound Extract or Pill, and combined with Henbane, which prevents griping. It is not to be given in pregnancy, nor when gastric or intestinal inflammation is suspected. The Tincture is ordered in Mixtures.

Dose—2 to 8 grains = 0.13 to 0.52 gramme

Ph Ger maximum single dose, 0.3 gramme, maximum daily dose, 1 gramme

Official Preparations—Extractum Colocynthidis Compositum and Pilula Colocynthidis Composita. Pilula Colocynthidis Composita is used in the preparation of Pilula Colocynthidis et Hyoscyami.

Not Official—Pilulæ Catharticæ Compositæ, Pilulæ Catharticæ Vegetabiles, Pilulæ Colocynthidis et Hydragyri, Tinctura Colocynthidis

— **Foreign Pharmacopœias**—Official in Austr., Belg., Dan., Dutch, Fr. (Coloquinte), Ger., Hung., Ital. (Coloquintida), Jap., Norw., Port. (Coloquintides), Russ., Mex. (Coloquintida), Swed., Swiss and U.S.

Descriptive Notes.—Colocynth fruits are imported from Asia Minor, Smyrna, Almeria in Spain, Mogador, Egypt and Cyprus, also from Persia, and more rarely from Marseilles and Trieste.

The kinds most frequently met with in commerce are the Turkish and Spanish, both of which have been peeled after drying, the Spanish less carefully, and shows more traces of the brown rind. The Mogador fruit is larger and unpeeled, it is brown externally when dry, though marbled with green and white when fresh, the Persian is peeled before drying, and consequently presents a shrunken appearance. It yields, however, the same proportion of pulp in relation to the seeds as the other varieties. Two forms of powdered Colocynth are sold, the one containing the seeds and the other freed from them as nearly as possible. The former is cheaper and is excluded by the Pharmacopœia direction 'freed from seeds,' and is used for keeping away moths from furs. The peeled fruit is alone official, occurring in more or less broken balls, about 2 in (5 cm) or less in diameter, *BP*, about 5 to 10 cm in diameter, *USP*. The Mogador variety exceeds the diameter given in *BP*, and is unpeeled and therefore excluded, whilst the Persian variety is not.

Tests—The boiled and cooled aqueous decoction of the pulp should yield no distinctive blue coloration with Starch Solution. The official test also adds that 'only traces of fixed oil should be removed by Ether.' The author has pointed out in the *Companion* (17th Edition) that the removal of the Seeds commercially is carried out very imperfectly, and as the Seeds contain about 15 p.c. of Oil it is doubtful whether a single trade sample could be found which would pass the official Ether Test, even on the supposition that the Pulp itself was free from Ether-soluble constituents. The Pulp, however, perfectly free from Seeds yields to Ether about 3 p.c. of extractive of an oily nature, so that the official test should be completely modified. Confirmation of the above results appears (*Analyst*, xxi, 31). Not merely Oil but also some gummy matter is extracted by Ether, both from the Seeds and the Pulp, and the use of a different solvent is therefore suggested. A limit for fixed Oil, as shown by extraction with Petroleum Spirit has been recommended, it should amount to from 1 to 2 p.c. The *BP* mentions that it yields when dried at 100° C (212° F) and incinerated, at least 9.0 p.c. of ash. The author has found the ash of the Pulp to vary between 8.6 and 14 p.c., and that of the Seeds between 2.2 and 4.0 p.c., on these figures Colocynth Pulp with an allowable 10 p.c. of Seeds would yield not less than 8 p.c. of ash. It should also be noted that the ash both of Pulp and Seed is very deliquescent. The figures given (*Analyst*, xxi, 31) for the Pulp are from 7.8 to 12.1 p.c., for the Seeds 1.7 to 3.2 p.c. The hygroscopic nature of the ash is commented on and a determination of the sulphated ash suggested. A limit of 9.0 to 12.0 p.c. of ash has been recommended. No ash limit is suggested in either the *USP* or *P.G.*

Preparations

EXTRACTUM COLOCYNTHIDIS COMPOSITUM COMPOUND
EXTRACT OF COLOCYNTH

Colocynth Pulp, 6, Extract of Barbados Aloes, 12, Scammony Resin, 4, Curd Soap, in shavings, 4, Cardamom Seeds, in the finest powder, 1, Alcohol (60 p c), 160

B P directs the Colocynth to be macerated in the Alcohol for four days, press out the Tincture, remove the Alcohol by distillation, and add the Extract of Aloes, Scammony Resin and Soap, evaporate to a firm extract, adding the Cardamoms towards the end of the process, but it is better to evaporate the Colocynth Extract to dryness, powder it, and mix with the other ingredients to form Pulv Ext Coloc Co, the product weighs about 24

6 of Compound Extract is about equal to $1\frac{1}{2}$ of Pulp (Simple Extract $\frac{3}{4}$), Extract of Aloes 3, Resin of Scammony 1, Curd Soap 1, Cardamoms $\frac{1}{2}$, Water $\frac{1}{2}$

Dose —2 to 8 grains = 0 13 to 0 52 gramme

Ph Ger maximum single dose of the Simple Extract, 0 05 gramme, maximum daily dose, 0 15 gramme

Commonly prescribed with Extract of Hyoscyamus, to prevent griping

Foreign Pharmacopœias —Official in Port, Colocynth 30, Aloes 55, Scammony 22, Hard Soap 15, Cardamoms 3, Swed, Colocynth 5, Aloes 10, Resin of Jalap 3, Cardamoms 1, Soap 2, Russ, Extract Colocynth 3, Aloes 10, Scammony 8, Extract of Rhubarb 5, U S, Extract Colocynth 16, Purified Aloes 50, Resin Scammony 14, Cardamoms 6, Soap 14, all alcoholic Not in the others Austr, Belg, Dan, Dutch, Ger, Hung, Ital, Jap, Mex, Port, Russ, Swiss and U S have a Simple Extract made with Alcohol

PILULA COLOCYNTHIDIS COMPOSITA. COMPOUND PILL OF
COLOCYNTH

Colocynth Pulp, 1, Barbados Aloes, 2, Scammony Resin, 2, Potassium Sulphate, $\frac{1}{4}$, Oil of Cloves, $\frac{1}{4}$, Distilled Water, *q s*
(about $\frac{1}{4}$) (about 1 in 6)

B P Dose —4 to 8 grains = 0 26 to 0 52 gramme

The *minimum* dose is somewhat high, as it is frequently prescribed in smaller doses The same may be said of the next pill, which is only two-thirds of the strength

For dispensing, keep the powders and oil ready mixed, and make up the mass as required with Water, or better still with Alcohol (60 p c)

This mass, when made with Scammony instead of Scammony Resin and divided into 5 grain pills, forms Gregory's pill

Foreign Pharmacopœias —Official in Norw, Colocynth 2, Aloes 4, Resin of Scammony 4, Oil of Cloves $\frac{3}{4}$, Suet 3, Glycerin 3, Swed, Compound Extract of Colocynth 7, Cloves 1, Resin of Jalap 2, Extract of Wormwood *q s*. Not in the others

PILULA COLOCYNTHIDIS ET HYOSCYAMI PILL OF
COLOCYNTH AND HYOSCYAMUS

Compound Pill of Colocynth, 2, Extract of Hyoscyamus, 1

B P Dose —4 to 8 grains = 0 26 to 0 52 gramme

Christison's Pill is 2 grains of *Pilula Colocynthidis et Hyoscyami* (*B P* 1867)

Hamilton's Pill —For some years past it has been the general practice to supply 4 or 5 grains of the *B P* pill mass, but some few houses in Edinburgh still supply the pills of the late Dr Hamilton, Jun, the formula for which was

Compound Extract of Colocynth, 2, Extract of Hyoscyamus, 1, mix and make into $\frac{1}{2}$ grain pills.

Foreign Pharmacopœias—Official in Jap, Colocynth 10, Aloes 20, Root of Jalap 20, Extract of Hyoscyamus 25, Potassium Sulphate 3, Oil of Cloves 1 Not in the others

Not Official

PILULÆ CATHARTICÆ COMPOSITÆ—Compound Extract of Colocynth, 16 grains, Mild Mercurous Chloride, 12 grains, Resin of Jalap, 4 grains, Gamboge, 3 grains, made into a mass with Diluted Alcohol (49 per cent) and divided into 12 pills—*U S P*

B.P.C. gives the same formula as above except that the *B.P.* Compound Extract of Colocynth is used in place of that of *U S P*, which is different

PILULÆ CATHARTICÆ VEGETABILES—Compound Extract of Colocynth, 12 grains, Extract of Hyoscyamus, 6 grains, Resin of Jalap, 4 grains, Extract of Leptandra, 3 grains, Resin of Peppermint, 2 minims, made into a mass with Diluted Alcohol (49 per cent), and divide into 12 pills—*U S P*

PILULÆ COLOCYNTHIDIS ET HYDRARGYRI *Syn* Abernethy's Pill.—Mercury Pill, 3 grains, Compound Extract of Colocynth, 2 grains, in one pill—*Pharm. Lond.*

This appears also in *B.P.C.*, but the proportions are reversed

TINCTURA COLOCYNTHIDIS—1 of Colocynth Pulp, in coarse powder, macerated with 10 of Alcohol (90 per cent) (1 in 10)

Dose—10 to 15 minims = 0.6 to 0.9 c.c. three times a day

Ph. Ger. maximum single dose, 1 gramme, maximum daily dose, 3.0 grammes

This has been incorporated in the *B.P.C.*

Foreign Pharmacopœias—Official in Hung and Mex, 1 in 5, Belg, Jap and Swiss, 1 in 10, Ger and Ital, Fruits 1, Alcohol 10, Swed, 1 in 10 with Anise Fruits $\frac{1}{10}$ Not in the others

Not Official

CONDURANGO CORTEX.

The Bark obtained from *Gonolobus condurango*

Medicinal Properties—It was introduced as a remedy for cancer, but it has not fulfilled the expectations formed of it. It relieves catarrh and hyperæsthesia of the stomach, and has been used with benefit in ulcer and cancer of the stomach, relieving the vomiting, pain and hæmatemesis, and improving the appetite—*J. M. R.* '82, 337, *L.* '95, 1004

Descriptive Notes—Formerly this bark was referred to *Gonolobus Condurango*, Truana, but in the *P.G.* it is now referred to *Marsdenia Condurango*, Reich, an Asclepiadaceous plant of Equatorial South America. It occurs in short gnarled pieces about 2 to 3 inches (50 to 75 mm) long or more, $\frac{1}{4}$ to $\frac{1}{2}$ inch (12 to 20 mm) broad and $\frac{1}{8}$ inch (3 mm) thick. Externally it is brownish grey, and the surface is uneven with slightly raised longitudinal ridges, the fracture is fibrous and exhibits numerous small glandular masses of white cells, but towards the outer surface a few projecting pores are perceptible. The distinctive features under the microscope are the large polygonal cells of the phelloderm cells, the abundance of spheraphid in the pith cells, the very variable sclerenchymatous cells, and the abundance of starch. The medullary rays consist of a single row of cells as seen in transverse section, and laticiferous vessels are present.

A cold infusion of the bark (1-2) becomes cloudy when heated, but becomes clear again when cold (*P.G.*)

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr., Ger, Jap, Mex., Norw, Russ, Span, Swed and Swiss. Not in the others.

EXTRACTUM CONDURANGO LIQUIDUM—A 1 in 1 fluid extract of Condurango bark prepared by percolation, with Alcohol (60 p c). The residue obtained after distilling the Alcohol from the latter portions of the percolate being dissolved in the reserved portion—*B P C Formulary* 1901, incorporated in the *B P C*

Dose—10 to 60 minims = 0.6 to 3.6 c c

100 of Condurango Bark is moistened with a mixture containing Alcohol (90 p c), 15, Distilled Water, 25, Glycerin, 10, and percolated with a mixture of Alcohol, 1, Distilled Water, 3, proceed as directed for fluid extracts, so as to produce 100—*Dan., Ger. and Jap.*

Austrian, Swedish, and Russian are similar, Dutch, Fluid extract, the bark is moistened with a mixture of Alcohol (90 p c) 60, Water 30, and Glycerin 10, and percolated with Alcohol (90 p c) 35 and Water 65, Belg, Fluid extract with Alcohol (30 p c), Solid extract with Alcohol (60 p c), Fr Fluid Extract with Alcohol (45 p c), Swiss, Fluid extract with a mixture of Alcohol (90 p c) 1 and Water 3, Spanish has an extract with Alcohol (70 p c)

VINUM CONDURANGO—Fluid Extract of Condurango, 1, Malaga Wine, 9—*Austr. and Swiss*

Condurango Bark, 1, Malaga, 10—*Dutch*

Condurango Bark, 1, Sherry, 10—*Ger. and Jap.*

Condurango Bark, 3, Alcohol (60 p c), 3, Carinena or Alicante Wine, *q s* to yield 100—*Span.*

All by weight

Fluid Extract of Condurango, 1, Detannated Sherry, *q s* to make 10—*B P C*

Dose—2 to 8 fl drms = 7.1 to 28.4 c c

CONIUM.

CONIUM

FR CIGUE, GER, SCHIERLING, ITAL, CICUTA, SPAN, CICUTA

The fresh Leaves and young Branches of *Conium maculatum*, L., as well as the dried unripe Fruits, are official in the *B P*, the full-grown but unripe Fruit, carefully dried and preserved, is official in the *USP*, the dried Leaves and flowering Stem Tops are official in the *PG*. The *USP* requires the Fruit to yield not less than 0.5 p c of Conium

Medicinal Properties—Sedative and antispasmodic, allays the cough in bronchitic affections, pertussis, and phthisis. Has been recommended in chorea and other spasmodic affections, also in visceral neuralgias and gastric pains. Applied externally in the form of ointment to ease pain of anal fissure or of hæmorrhoids, and cancer

Dose—Of the Succus 1 to 2 fl drms = 3.6 to 7.1 c c. Of the Tincture 30 to 60 minims = 1.8 to 3.6 c c

Ph. Ger. maximum single dose, 0.2 gramme, maximum daily dose, 0.6 gramme

Prescribing Notes—In consequence of the great variation in strength of Conium preparations, the standardised Fluid Extract or Conium Hydrobromidum should be prescribed. 1 fl drms of the Fluid Extract is about equal to 1 fl oz of Succus Conii (average strength)

Incompatibles.—Caustic Alkalis, and vegetable Astringents.

Official Preparations—Succus Conii from the Folia Unguentum Conii from the Succus Tinctura Conii from the Fructus

Not Official—Extractum Conii, Extractum Conii Liquidum, Pessus Coniinae, Pilulae Conii Compositae, Vapor Coniinae, Coniina, Coniinae Hydrobromidum, and Coniinae Hydrochloridum

Antidotes—In case of poisoning by Hemlock, stomach-tube or emetics, followed by 10 minims of Strychnine hypodermically, artificial respiration

CONII FOLIA. CONIUM LEAVES.

The fresh Leaves and younger Branches of *Conium maculatum*, collected when the fruit begins to form

Foreign Pharmacopœias—Official in Austr, Ger, Mex, Port and Span Not in Belg, Dan, Dutch, Fr, Hung, Ital, Jap, Norw, Russ, Swed, Swiss or U S

Descriptive Notes.—Conium has pinnately decompound leaves and is distinguished from other British umbelliferous plants having similar leaves by the purplish spots on the stem and petioles of the leaves, by the mouse-like odour evolved when rubbed with Liquor Potassae, and by the leaves being quite free from hairs. The extreme points of the leaf segments are white or colourless, whilst in *Aethusa Cynapium*, L., which has also hairless leaves, the tips are brown, and it has no general involucre to the umbels as in Conium, but only a partial one of three long bracts. The fresh leaves of Conium only are official and are in best condition at the end of June and beginning of July. Such of the British species of the genus *Cherophyllum* as resemble Conium in appearance, have hairy leaves and cylindrical fruits. Under the microscope it is distinguished by striated epidermal cells, by the parenchymatous cells containing minute, usually single, crystals of Calcium Oxalate, the absence of hairs, and the presence of annular as well as spiral vessels.

Tests.—Conium leaves when bruised possess a strong and peculiar odour, somewhat resembling that of mice, and when the leaves are rubbed with Potassium or Sodium Hydroxide Solution the odour is intensified.

Preparations

SUCCUS CONII. JUICE OF CONIUM

3 of Juice, obtained from the fresh Leaves and young Branches, preserved by the addition of 1 of Alcohol (90 p c)

Dose.—1 to 2 fl. drm = 3.6 to 7.1 cc

Much larger doses are also given

UNGUENTUM CONII. CONIUM OINTMENT

Evaporate 8 of the Juice of Conium on a water-bath to 1, at a temperature not exceeding 110° F (60° C), and mix with 3 of Hydrous Wool Fat

Contrary to what might have been expected, the alkaloidal strength of the juice is not affected by the evaporation, but it is better to evaporate the juice to 2 and to use Anhydrous Wool Fat, also to add 2 p c of Boric Acid

Becomes mouldy on keeping—P J '98, 165, 232

Not Official

EXTRACTUM CONII —Made from the fresh leaves and young branches of Hemlock —*B P* 1885

This has been incorporated in the *B P C*

PILULÆ CONII COMPOSITÆ —Extract of Hemlock, 5, Ipecacuanha, in powder, 1, Treacle *q s* —*B P* 1885

This has been incorporated in the *B P C*

VAPOR CONIINÆ —Juice of Hemlock, $\frac{1}{2}$ fl oz, Solution of Potash, 1 fl dr, Distilled Water, 1 fl oz —*B P* 1885

This has been incorporated in the *B P C*

CONII FRUCTUS CONIUM FRUIT

The dried, full-grown, unripe Fruits of *Conium maculatum*

Conium Fruits are not officially required to yield any definite percentage of Coniine. The *U S P* states that they shall yield not less than 0.5 p.c. of Coniine. The standard adopted by the *U S P* has been criticised as being too low, but the standard is justified (*Y B P* '05, 398)

Foreign Pharmacopœias —Official in Fr, Mex, Port, Span and U S

Descriptive Notes —Conium fruits are ovoid, greyish-green, slightly compressed laterally, and, as met with in commerce, consist of the separate mericarps. The size is given in *B P* as $\frac{1}{8}$ inch (3 mm) long, and nearly as broad, in *U S P* as 3 mm long, and about 1.5 mm in diameter. The five dorsal ridges are more or less wavy and irregularly crenate, this feature being most conspicuous before the fruits are fully ripe. The surface between the ridges is glabrous but minutely wrinkled. The flat surface shows a narrow deep depression which gives a reniform outline to a transverse section of the mericarp. The *U S P* states that Conium fruit after being kept for more than two years is unfit for use. Although vittæ are present in the very young fruit they subsequently disappear and are absent in the fruit when mature. Conium Fruits are characterised under the microscope by thin-walled nearly cubical cells, which form a layer outside the endosperm. The Coniine and an essential oil are contained in the cells of the endocarp, so that a finely powdered fruit is not necessary for its extraction.

Tests —The percentage of Coniine present in Conium Fruits may be determined by extraction with suitable solvents, and weighing the alkaloid as a Hydrochloride. A weighed quantity of 5 grammes of the finely-powdered fruit is extracted with 50 c.c. of a saturated solution of dry Hydrochloric Acid gas in Chloroform. The extraction with a further quantity of a similar mixture is continued until 6 drops of the chloroformic mixture evaporated on a watch-glass, and the residue acidified with Diluted Sulphuric Acid, gives no precipitate with Mayer's reagent. The mixed chloroformic liquids after separation from the marc are shaken with two separate quantities of 25 c.c. each of Water. The mixed aqueous shakings are in turn separated, shaken twice with 10 c.c. of Chloroform, and the Chloroform separated. The aqueous portion is made alkaline by the addition of Sodium Hydroxide Solution, and the liberated alkaloid extracted by shaking with three

successive portions each of 10 c c of Chloroform. The chloroformic liquids are separated in each case, mixed, run into 10 c c of the saturated solution of dry Hydrochloric Acid gas in Chloroform, evaporated to dryness on a water-bath, the residue dried at a temperature not exceeding 90° C (194° F), and weighed, 162.41 parts of anhydrous Coniine Hydrochloride represent 126.22 parts of Coniine.

The method of determination adopted by the *U S P* is essentially as follows.—A weighed quantity of 10 grammes of the Fruit in No. 60 powder is shaken at intervals during four hours in an Erlenmeyer flask, with 100 c c of a mixture of 98 parts of Ether, 8 parts of Alcohol (94.9 p c), and 3 parts by volume of Ammonia Water. A measured quantity of 50 c c of the clear liquid is decanted into a beaker, and mixed with sufficient Normal Volumetric Sulphuric Acid Solution to produce an acid reaction. The Ether is evaporated on a water-bath, 15 c c of Alcohol (94.9 p c) added, and the mixture set aside for two hours to permit of the Ammonium Sulphate depositing, the Tincture filtered, the residue and filter washed with a little Alcohol (94.9 p c), and the washings mixed with the filtrate. The excess of acid is neutralised by Sodium Carbonate, a slight acidity being carefully maintained. The liquid is now carefully concentrated on a water-bath to a volume of 3 c c, mixed with an equal volume of Water and 2 drops of Normal Volumetric Sulphuric Acid Solution. The liquid is washed with two successive quantities each of 15 c c of Ether, the ethereal liquids separated, the acid liquid transferred to a separator, sufficient Sodium Carbonate Test-solution added to render the liquid distinctly alkaline to red Litmus paper, and the liberated alkaloid shaken out with successive portions of 15 c c, 15 c c, and 10 c c of Ether. The ethereal solutions are in each case separated, transferred to a tared beaker, and sufficient 5 p c Hydrochloric Acid solution added to the mixed ethereal solutions to render them distinctly acid. The Ether is removed by evaporation at a gentle heat on a water-bath, and the excess of Hydrochloric Acid by adding two separate quantities of 3 c c each of Alcohol (94.9 p c), and removing them in each case by evaporation. The residue is dried at a temperature not exceeding 60° C (140° F). This weight, multiplied by 0.777 and the product by 20, gives the percentage of Coniine present in the Fruit.

Coniine may be titrated with Normal or Deci-normal Volumetric Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutrality, 1 c c of the Normal Acid represents 0.012622 gramme of Coniine.

Preparation.

TINCTURA CONII. TINCTURE OF CONIUM

1 of Conium Fruit, recently reduced to No. 40 powder, percolated with Alcohol (70 p c), *qs* to yield 5.
Now 1 in 5 instead of 1 in 8.

Dose.—30 to 60 minims = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Port, Tinct Cicutæ, 1 in 5, also Fresh Herb 1, Spirit 1, Mex, Leaves, 1 in 5 Not in the others

Tests—Tincture of Conium possesses a specific gravity of from 0.895 to 0.900, contains about 1.75 p.c. of total solids and about 68 p.c. w/v of Absolute Alcohol. The *B.P.* Tincture is not a standardised preparation, and no process for the quantitative determination of the Conium is given.

Not Official.

EXTRACTUM CONII LIQUIDUM—A standardised Liquid Extract prepared by treating 100 of Conium Fruit (in No. 40 powder) with a mixture of 100 of Alcohol (60 p.c.) and $1\frac{1}{2}$ of Acetic Acid, the exhaustion is completed with Alcohol (60 p.c.), finally the liquid is adjusted to contain 1 p.c. w/v alkaloidal hydrochlorides—*B.P.C. Formulary* 1901, incorporated in the *B.P.C.*

Dose—5 to 15 minims = 0.3 to 0.9 c.c.

The *U.S.P.* has also a Fluidextractum Conii, about 1 in 1, which is standardised to contain 0.45 p.c. w/v of Conium. It is prepared by treating 100 of Conium Fruit in No. 40 powder with a mixture of 98 of Alcohol (60 p.c.) and 2 of Acetic Acid (*U.S.P.*), the exhaustion is completed with Alcohol (60 p.c.)

Tests—The *U.S.P.* mixes a measured quantity of 10 c.c. of the Fluid Extract with a little clean sand and evaporates to dryness on a water bath. After the Extract and the sand are uniformly mixed they are transferred to an Erlenmeyer flask and shaken at intervals during one hour with 100 c.c. of a mixture of 100 c.c. of Ether, 7 c.c. of Alcohol (94.9), and 3 c.c. of Ammonia Solution, the dish being washed out with the mixture, and the mixture added in portions. A measured quantity of 50 c.c. of the clear liquid is decanted into a beaker, and sufficient Normal Volumetric Sulphuric Acid Solution added to produce a distinctly acid reaction. The Ether is removed by evaporation, 15 c.c. of Absolute Alcohol added, and the Ammonium Sulphate allowed to deposit during two hours. The liquid is filtered, the residue and filter washed with a little Absolute Alcohol, the washings being added to the filtrate. The excess of acid is neutralised by the careful addition of Sodium Carbonate Test Solution, a slight acidity being maintained. The liquid is concentrated by evaporation on a water bath, to a volume of 8 c.c., mixed with an equal volume of Water and 2 drops of Normal Volumetric Sulphuric Acid Solution. The acid liquid is washed with two successive portions each of 15 c.c. of Ether, the acid liquid separated in each instance, it is transferred to a separator, sufficient Sodium Carbonate Test Solution added to render the liquid distinctly alkaline to red Litmus paper, and the liberated alkaloid removed by extraction with successive portions of 15 c.c., 10 c.c., and 10 c.c. of Ether. The ethereal solutions are separated in each case, transferred to a tared beaker, mixed, and sufficient of a 5 p.c. Hydrochloric Acid Solution added drop by drop to ensure an excess of acid. The Ether is then removed by distillation, the excess of Hydrochloric Acid by evaporating twice with successive portions of 3 c.c. of Alcohol (94.9 p.c.), the residue dried at a temperature not exceeding 60° C (140° F), till constant in weight, and weighed after cooling in a desiccator. The weight multiplied by 0.777 and the product by 20 gives the p.c. w/v of Conium present in the sample of Fluid extract.

The *B.P.C.* states, 'determine the proportion of alkaloids in the strong liquid extract, and adjust the finished product so that it shall contain alkaloids equivalent to 1.0 p.c. of alkaloidal hydrochlorides,' but does not give the details of any process by which they may be determined.

PESSUS CONIINÆ—Conium, $\frac{1}{2}$ maxim, Gelatin Basis, 20 grains—*Women*

Conium Hydrobromide, $\frac{1}{2}$ grain, Oil of Theobroma, 120 grains—*B.P.C.*

CONIINA. *Syn.* Cicutine $C_8H_{17}N$, eq. 126.22—A colourless, or pale yellow, volatile oily liquid, with a characteristic penetrating mousey odour. Obtained

from *Conium maculatum*. It unites with acids to form crystalline salts, which are much more stable than the alkaloid.

Solubility—1 in 100 of Water. It mixes in all proportions with Alcohol (90 p.c.) and with Ether.

Causes a very great increase in the blood pressure when injected (*L* '05, 1851), the effect, however, is transient, and prolonged administration causes paralysis.

Dose—It has been given in doses of $\frac{1}{12}$ grain to 1 grain = 0.0054 to 0.06 gramme, but the Foreign Pharmacopœias give much smaller doses, 1 to 4 milligrammes = $\frac{1}{24}$ to $\frac{1}{18}$ grain.

Foreign Pharmacopœias—Official in Mex. Not in the others.

Tests—Conine has a specific gravity of 0.886 (Schorm), 0.844 (Ladenburg). It boils at about 169° C (336 2° F). It is dextrogyrate, its specific rotation being +13.8° for the Sodium ray.

The aqueous solution of Conine is powerfully alkaline in reaction. When a glass tube with concentrated Hydrochloric Acid is held closely over a small quantity of Conine contained in a watch-glass, white fumes are produced, and if sufficient of the fumes be passed over the surface of the alkaloid, it is wholly converted into a crystalline Hydrochloride, Nicotine Hydrochloride is amorphous.

On the addition of a large excess of concentrated Hydrochloric Acid to Conine a pale red tint is produced, gradually deepening in colour, Sulphuric Acid gives no immediate change with pure Conine, but the mixture gradually becomes purple-red and then olive-green. It may be distinguished from Nicotine by producing with Mercuric Chloride Solution a white amorphous, instead of a crystalline precipitate, by its non-precipitation with Platonic Chloride Solution, and by its alkalinity. In Phenolphthalein Solution, an aqueous solution of Conine being coloured red immediately on the addition of 1 or 2 drops of Phenolphthalein Solution, Nicotine is neutral to Phenolphthalein Solution. When heated slowly with free access of air it is completely volatilised, leaving no weighable residue.

CONIINÆ HYDROBROMIDUM Conine Hydrobromide $C_8H_{11}N.HBr$, eq. 206.57—Transparent, colourless, rhombic crystals, or a white, crystalline powder. The usual form for prescribing Conine, of which it contains about 60 p.c.

It should be protected as far as possible from the light in well-closed glass bottles of a dark amber tint.

Solubility—1 in 2 of Water, 1 in 3 of Alcohol (90 p.c.)

Dose— $\frac{1}{2}$ to 2 grains = 0.01 to 0.13 gramme.

For hypodermic use, $\frac{1}{12}$ grain in 5 minims of Water.

Official in Fr. (1908) and Mex.

Tests—Conine Hydrobromide melts according to *Fr. Codex* (1908) at 211° C (411.8° F). The Conine obtained from the Hydrobromide answers the tests distinctive of the alkaloid given under that heading. When moistened with a few drops of Potassium or Sodium Hydroxide Solution, the peculiar mousy odour of Conine is evolved. The aqueous solution acidified with diluted Nitric Acid gives on the addition of Silver Nitrate Solution a yellowish-white curdy precipitate, practically insoluble in Ammonia Solution and in Nitric Acid. The salt should leave no weighable residue when ignited with free access of air.

CONIINÆ HYDROCHLORIDUM—Colourless crystals, readily soluble in Water and in Alcohol (90 p.c.).

Tests—The Conine obtained from Conine Hydrochloride responds to the test for the alkaloid given under that heading. When moistened with 1 or 2 drops of Potassium or Sodium Hydroxide Solution, the peculiar mousy odour of Conine is evolved. The aqueous solution acidified with Nitric Acid yields with Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, soluble in Ammonia Solution. The salt when incinerated with free access of air leaves no weighable residue.

Not Official

CONVALLARIA

The entire Plant of *Convallaria Mayalis*, L (Lily of the Valley), gathered when the flowers commence to open and dried

Medicinal Properties—A cardiac tonic, diuretic Not cumulative like Digitalis, but according to Mitchell Bruce it is a very uncertain remedy It has been long employed by the Russian peasantry as a remedy for dropsy The late Professor Sée considered that it may be used in all forms of heart failure, for it has none of the nauseating effects of Digitalis, nor does it exhaust the contractility of the heart and arteries

The juice of fresh plant stated (*P J* '04, 11 967) to contain 0.225 p.c. Convallamarin, and 0.12 p.c. Convallarin

Foreign Pharmacopœias—Official in Austr, Fr (Muguet), Ital (Mughetto), Mex, Span (Lirio de los Valles) and U.S. Not in the others

Convallaria contains 2 glucosides—Convallarin, a purgative, and Convallamarin, allied to Digitalin in its action on the heart, the dose of the latter is $\frac{1}{8}$ to 2 grains = 0.008 to 0.13 gramme

EXTRACTUM CONVALLARIÆ (*Fr, Ital and Span*)—An aqueous extract of the Stalks and flowers of Convallaria freshly gathered and dried Mex, from roots

Dose—2 to 5 grains = 0.13 to 0.32 gramme three times a day

Ital maximum single dose, 0.20 gramme, maximum daily dose, 1.0 gramme

FLUIDEXTRACTUM CONVALLARIÆ (*U.S.*)—1 in 1, from the rhizome and roots of Convallaria, with a mixture of Alcohol (95 p.c.) 650, and Water 350 = about Alcohol (60 p.c.)

Average Dose—8 minims = 0.5 c.c.

Russ has a Tincture from fresh Flowers

B.P.C. has 1 in 1 Fluid Extract, dose 5 to 10 minims, and a Tincture 1 in 8, dose 5 to 20 minims, both are made from the dried Flowers with Alcohol (60 p.c.)

COPAIBA.

COPAIBA

B.P.Syn—COPAIBA

FR, COPAHU, GER, COPAIVABALSAM, ITAL, BALSAMO DI COPAIBA, SPAN, OLEO RESINA DE COPAIBA

An Oleo-Resin, obtained from the trunk of *Copaifera Lansdorfii*, Desf, as well as from other species of *Copaifera*

Obtained from the northern part of South America The commercial varieties Para, Maianham, Maracabo, and Angostura are named from the various ports of shipment

Solubility—(Nearly clear) 1 in 1 (*or less*) of Alcohol (90 p.c.), but if more Alcohol be added it becomes cloudy, in all proportions of Absolute Alcohol, Ether, Benzol, and the fixed and volatile Oils, also in four times (*or less*) its bulk of Petroleum Spirit, the solution only yielding a filmy deposit on standing, also 1 in 2 (*or less*) of Glacial Acetic Acid

Medicinal Properties—Stimulant, antiseptic, and diuretic Acts more particularly upon the mucous membrane of the genito-urinary tract Used in gonorrhœa, after the acute stage has passed, and in gleet Sometimes combined with Buchu and Cubebs

Useful in chronic bronchitis and bronchiectasis, when a disinfectant expectorant is indicated. The resin is used as a diuretic in cardiac and hepatic dropsy, but not in renal, as it is liable to irritate the kidneys.

Dose.—30 to 60 minims = 1 8 to 3 6 c c

Prescribing Notes.—*Can be given in the form of pills or paste (see below), also in capsules. It may be suspended in Water by means of Mucilage of Gum Acacia (see p 3), or Liquor Potassæ, which saponifies it. Cinnamon Water, Peppermint Water, the Tinctures of Orange and Ginger have been used as flavouring agents. The Oil of Copaiba can be suspended by means of Mucilage, as can also the Resin of Copaiba.*

When Copaiba is boiled with Solution of Potassium Hydroxide the Oil is destroyed, and the Resin separates on standing. The liquid portion is consequently miscible with Water.

Official Preparation.—Oleum Copaibæ

Not Official.—Electuaire de Copahu Composé, Liquor Copaibæ Solubilis, Liquor Copaibæ, Buchu et Cubebæ, Liquor Copaibæ cum Santalo, Liquor Copaibæ et Buchu, et Cubebæ cum Santalo, Haustus Copaibæ, Mistura Copaibæ, Mistura Copaibæ Acidæ, Mistura Copaibæ Alkalina, Pasta Copaibæ, Pilula Copaibæ, Resina Copaibæ

Foreign Pharmacopœias.—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port (Terebinthina Copahiba), Russ, Span, Swed, Swiss and U S

Descriptive Notes.—The oleo-resin, incorrectly designated in commerce Balsam of Copaiba, varies considerably in consistence, according to the species of Copaifera from which it is derived. These, besides the species mentioned in the *B P*, viz, *C Lansdorffii* (the source of Maranham Copaiba), are believed to be *C officinalis*, L (Maracaibo and Cartagena Copaiba), *C Guianensis*, Desf (Surinam Copaiba and B Guiana Copaiba), *C coriacea*, Mart (Bahia Copaiba), *C confertiflora*, Benth, *C oblongifolia*, Mart, and *C rigida*, Benth, are also believed to yield some of the Copaiba of commerce. The *B P* lays no restriction upon the species from which it may be derived. The Para kind (*C multijuga*, Hayne), which is the most fluid and the palest in colour, almost resembling Olive oil in colour and consistence, yields 60 to 90 p c of volatile oil, and is therefore chiefly used for distilling the oil. The Maracaibo, which is thicker and of a brownish colour, like that of Maranham, and with a greenish fluorescence, yields only about 40 p c, and that from Bahia about 50 p c, the Maranham kind is preferred in Germany.

When a Copaiba containing much oil is desired, the Para, Surinam, and British Guiana kinds are most suitable, and when one containing much resin is required, the Maranham and Maracaibo and Cartagena are preferable. Gurgun balsam resembles the darker varieties of Copaiba in colour and consistence, but if heated to 130° C (266° F) it is transformed into a jelly.

Tests.—Copaiba has a specific gravity of from 0 985 to 0 994, *B P* gives 0 916 to 0 993, the *USP* 0 950 to 0 995 at 25° C (77° F), the *PG* 0 980 to 0 990. It should contain at least 40 p c of volatile oil, which should possess an optical rotation of from -14° to -17°, and a boiling point of 245° to 275° C.

(473° to 527° F), *B.P.*, not under 250° C (482° F) This volatile oil should not possess an odour of Turpentine, and if after its volatilisation the Balsam be heated until the volatile constituents are removed a hard readily friable residue should remain The Acid and Ester value afford a useful criterion of the quality of the Balsam, but a method for their determination is not included in the *B.P.* The *USP* and the *PG* include a determination of the Acid value, the *USP* requiring from 64.10 to 89.18, the *PG* 75.8 to 84.24 The *PG* includes a determination of the Ester value, which should be not less than 8.2, but no ester value is included in the *USP* The methods of determination given in the *PG* have been severely criticised, and are considered to be of little value, the only variety of Balsam which at all approximates the figures given being the Maranhão variety The figures for the optical rotation of Copaiba given in the *B.P.* require to be halved (*CD* '00, 1166; '03, 118; *PJ* '00, 154; '00, 1198), as they represent the rotation of the plane of a ray of polarised light in a tube of 200 mm., and not in a tube of 100 mm. as adopted in other cases It is probable that even when halved the limits may be too narrow, as some genuine samples of Copaiba yield an oil having a rotation as low as -9°, and Gildemeister and Hoffmann quote -7° to -35° as the rotation of Copaiba Oil from Pará or Maracaibo Balsam It has been suggested (*CD* '00, 1166, *PJ* '00, 1199) that the *B.P.* might with advantage include a method of determining the percentage of volatile oil in the Balsam, such as by heating a weighed quantity of the Balsam on a water-bath, and finally in a water-oven at a temperature of just below 100° C (212° F), until constant in weight Determinations of the Acid and Ester values of ten samples of the Balsam showed them, with one exception, to vary between 10.64 and 19.60 for the Ester, and 78.96 and 93.52 for the Acid value

It has also been pointed out (*CD* '01, 1436, *PJ* '01, 1326) that it would be useful to include monographs for the Resin as well as the Volatile Oil The Volatile Oil might be required to possess a specific gravity of 0.903 to 0.908, an optical rotation in a 100 mm. tube of -7° to -21°, a boiling point of 245° to 275° C (473° to 527° F), and a solubility in Absolute Alcohol of 1 in 1 The Resin should be soluble in Alcohol (90 p.c.), Ether, and Carbon Disulphide, and should possess an Acid value of at least 119.7 The specific gravity suggested for the Balsam is 0.97 to 0.995, and the Acid Value for the Balsam of at least 75.20

The Acid value of Maracaibo Copaiba Balsam should be between 76.52 and 94.90, the Ester value from 0.47 to 8.75, the Acid value of Pará Balsam should be from 65.8 to 72.0, and the Ester value from 1.9 to 2.9

The more generally occurring adulterants of Copaiba Balsam are Turpentine oil, fixed oils, *eg.*, Olive and Castor Oils, Paraffin Oils, Gurjun Balsam, Colophony, and African Copaiba Balsam Turpentine Oil may be readily detected by the odour on evaporation and the boiling point and rotation of the volatile Oil Fixed oils are indicated by the character of the Resin remaining after the

volatilisation of the ethereal Oil, and by the solubility of the Balsam in Alcohol (90 p c), and in Petroleum Ether, the *USP* includes a test with 20 drops of the Balsam and 1 c c of a 1 in 10 alcoholic Potassium Hydroxide Solution, mixing when cool with twice the volume of Ether, no gelatinisation should occur. Paraffin Oils are also readily detected by the solubility of the sample in Alcohol (90 p c), when a measured quantity of 5 c c of the Balsam is shaken with 15 c c Alcohol (94.9 p c), the mixture boiled for one minute, allowed to cool and to stand for one hour, no drops of oil should separate. Gurjun Balsam may be detected by the Nitric and Sulphuric Acid, and Acetic and Nitric Acid tests, the *BP* employs a cooled mixture of equal parts of Nitric and Sulphuric Acids, and Glacial Acetic Acid containing a small quantity of Nitric Acid, as tests for the absence of Gurjun Balsam, no transient violet coloration should be produced when 1 drop of the former mixture is added to 2 drops of the Balsam dissolved in 20 parts of Carbon Bisulphide, nor should a reddish or purple colour be yielded when 4 drops of the Balsam are carefully added to $\frac{1}{2}$ oz of Glacial Acetic Acid, to which has been added 4 drops of Nitric Acid. The *USP* uses Glacial Acetic Acid mixed with a few drops of Nitric Acid in performing this test, but gives more explicit directions for its application. No reddish zone should be produced, nor should the fluid assume a red or purple colour when 4 drops of Copaiba are carefully poured on top of a mixture of 1 drop of Nitric Acid (sp gr 1.40) and 3 c c of Glacial Acetic Acid. Colophony may be detected by shaking 1 gramme of Copaiba, in a stoppered vial, with 10 c c of Ammonia Solution, when allowed to stand 24 hours it should not gelatinise, nor should a firm mass be produced, but the liquid will become turbid. African Copaiba yields on distillation a volatile oil which is dextrogyrate, and its presence may be ascertained by its effect on the optical rotation of the distilled oil.

Volumetric Determination—1 gramme of Copaiba, dissolved in 50 c c of Alcohol, should require not less than 2.3 c c and not more than 3.2 c c (2.7 c c to 3 c c, *PG*) of the Semi-normal Volumetric Alcoholic Solution of Potassium Hydroxide for neutralisation, using 1 c c Phenolphthalein Solution (10 drops, *PG*) as indicator, *PG* and *USP*. If a further addition of 20 c c of Semi-normal Volumetric Alcoholic solution of Potassium Hydroxide be made, and the mixture warmed for 15 minutes on a water-bath, and titrated with Semi-normal Volumetric Solution of Hydrochloric Acid, it should require for the neutralisation of the excess of Potassium Hydroxide at least 19.7 c c of the Acid Solution, *PG*.

OLEUM COPAIBÆ. OIL OF COPAIBA

A yellow, or yellowish-brown, oily liquid, distilled from Copaiba. It has a distinctive Copaiba odour and a bitter, persistent taste.

It should be kept in well-closed glass bottles of a dark amber tint in a cool atmosphere, and protected as far as possible from the air and light.

Copaiba Oil contains, according to Gildemeister and Hoffmann, a sesquiterpene Caryophyllene, yielding Caryophyllene Hydrate in crystals, melting at 96° C (204.8° F) on treatment with Glacial

Acetic Acid and Sulphuric Acid A crystalline acid melting at 140°C (284°F), which has been identified as a symmetric Dimethyl-succinic Acid, has also been noted, but doubt is expressed as to whether the acid owes its origin to Caryophyllene or some other minor constituent of the oil

Solubility—1 in 20 of Alcohol (90 p c), nearly insoluble in Alcohol (60 p c), mixes in all proportions with Absolute Alcohol

Dose—5 to 20 minims = 0.3 to 1.2 c c

Foreign Pharmacopœias—Official in U S Not in the others

Tests—Copaiba Oil has a specific gravity of 0.903 to 0.908, the gravity varying considerably with the age of the oil and its exposure to the air—the official figures are 0.900 to 0.910, it is laevogyrate, the optical rotation being from -7° to -21° in a 100 mm tube. It boils between 245° and 275°C (473° and 527°F). It is neutral in reaction, and is soluble in its own volume of Absolute Alcohol. The more generally occurring adulterants are African Copaiba Oil and Gurjun Balsam Oil. The former is dextrogyrate, and may be detected by its effect on the optical rotation of the oil, the latter is recognised by its higher specific gravity and greater optical activity, which amounts generally to about -35° to -130° in a 100 mm tube, though dextrogyrate oils are also known to exist. The presence of Gurjun Balsam Oil may also be detected by the test with Acetic and Nitric Acids given under Copaiba.

Not Official

ELECTUAIRE DE COPAHU COMPOSÉ—Copaiba, 100, Cubebs in powder, 150, Catechu in powder, 50, Oil of Peppermint, 3—*Fr*

LIQUOR COPAIBÆ SOLUBILIS—Boil 20 of Copaiba with 30 of Solution of Potash for an hour, add 10 of Water, and mix thoroughly. Set aside until cold and well separated, draw off the clear liquor from the upper only portion and the sediment, and evaporate it to 38, to this add 2 of Solution of Potash—*Pharm Form* and the *Australian Pharmaceutical Formulary*

This has been incorporated in the *B P C*

LIQUOR COPAIBÆ, BUCHU ET CUBEBÆ—Liquid Extract of Buchu, 1, Liquid Extract of Cubebs, 1, Solution of Copaiba, 8—*Pharm Form*

This has been incorporated in the *B P C*

LIQUOR COPAIBÆ CUM SANTALO—Oil of Santal, 1, Alcohol (90 p c), 1, Solution of Copaiba, 8. The liquor is sometimes flavoured with Cinnamon or other essential oil, 5 to 10 minims to the ounce. Oil of Sandal Wood can also be combined with Solution of Copaiba, Buchu et Cubebs in the same manner as above—*Pharm Form*

Solution of Copaiba, 80, Oil of Sandal Wood, 10, Oil of Cassia, $\frac{1}{2}$, Alcohol (90 p c), q s to make 100—*B P C*

LIQUOR COPAIBÆ ET BUCHU ET CUBEBÆ CUM SANTALO—Solution of Copaiba, Buchu, and Cubebs, 80, Oil of Sandal Wood, 10, Oil of Cassia, $\frac{1}{2}$, Alcohol (90 p c), q s to produce 100—*B P C*

HAUSTUS COPAIBÆ—Copaiba, 15 minims, Solution of Potassium Hydroxide, 5 minims, Spirit of Nitrous Ether, 15 minims, Mucilage of Gum Acacia, 60 minims, Camphor Water, to 1 fl oz—*St Bartholomew's*.

MISTURA COPAIBÆ (Lafayette)—Copaiba, 4, Spirit of Nitrous Ether, 4, Compound Tincture of Lavender, 4, Solution of Potassium Hydroxide, 1, Syrup, 10, Mucilage of Acacia, q s to make 32. Mix the Copaiba with the

Solution of Potassium Hydroxide and the Spirit of Nitrous Ether, then add the Compound Tincture of Lavender, and lastly the Syrup and Mucilage of Acacia Well mix by shaking—*U S N F*

MISTURA COPAIBÆ (*Chapman*)—Copaiba, 8, Spirit of Nitrous Ether, 8, Compound Tincture of Lavender, 2, Tincture of Opium, 1, Mucilage of Acacia, 4, Water, *q s* to make 32—*U S N F*

MISTURA COPAIBÆ—Copaiba, 15 minims, Mucilage of Acacia, 60 minims, Magnesium Sulphate, 30 grains, Cinnamon Water, to 1 fl oz—*St Mary's*

C L U O I, 20 minims, Tincture of Quillaia, 20 minims, Spirit of Nitrous Ether, 30 minims, Camphor Water, to 1 fl oz—*Channing Cross*

Copaiba, 15 minims, Mucilage of Gum Acacia, 30 minims, Water, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

MISTURA COPAIBÆ ACIDA—Copaiba, 20 minims, Dilute Sulphuric Acid, 10 minims, Mucilage of Acacia, $\frac{1}{2}$ fl oz, Water, to 1 fl oz—*King's*

MISTURA COPAIBÆ ALKALINA—Copaiba, 20 minims, Solution of Potash, 10 minims, Mucilage of Acacia, 2 fl dr, Water, to 1 fl oz—*King's*

PASTA COPAIBÆ—Copaiba, 8, Powdered Cubebs, 24, Extract of Hyoscyamus, 1, Camphor, 1, Treacle, *q s*

Dose—A piece the size of a filbert nut three or four times a day in gonorrhœa—*L. '88*, 1 1019

PILULA COPAIBÆ—Copaiba, 94, Magnesia, 6, mix intimately and set in a cold place. Should the mixture not concreate in eight or ten hours, the copaiba should be shaken with $\frac{1}{10}$ of its weight of Water, then the uncombined Water allowed to subside and the Copaiba poured off

Foreign Pharmacopœias—Official in Span (Pildoras de Copaiba)

RESINA COPAIBÆ—Prepared from the Oleo-resin by distilling off the Volatile Oil

A yellowish, or brownish-yellow, brittle resin, with an acid reaction Soluble in Alcohol

Tests—Copaiba Resin is soluble in Alcohol (90 p c), Ether, and Carbon Bisulphide It possesses an Acid value of not less than 119.77

CORIANDRI FRUCTUS.

CORIANDER FRUIT

FR., CORIANDRE, GER., KORIANDER, ITAL., CORIANDRO, SPAN., CILANTRO

The dried, ripe Fruit of *Coriandrum sativum*, L

Medicinal Properties—Stimulant, aromatic, and carminative

Dose.—20 to 60 grains = 1.3 to 4 grammes

Official Preparation—Oleum Coriandri Contained in Confectio Sennæ, Syrupus Rhei, Tinctura Rhei Composita, and Tinctura Sennæ Composita The Oil is contained in Syrupus Sennæ

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Hung., Mex (Culantro), Norw., Port (Coentro), Span and US Not in Ger., Ital., Jap., Russ., Swed or Swiss

Descriptive Notes.—The Coriander Fruit of commerce has the two mericarps united, and is globular, about $\frac{1}{4}$ inch (5 mm) in diameter, of a buff or light brown colour (brownish yellow, *B P*), glabrous, crowned with minute calycine teeth, and the conical base of the two slender divergent styles There are four prominent

secondary ridges, and five inconspicuous wavy primary ridges alternate with them on each mericarp, but there are no vittæ between them, there being only two vittæ on the concave commissural surface of each mericarp. The taste and odour of the dried fruit are agreeably aromatic and characteristic, but in the unripe fruit the odour is disagreeable. The fruits are imported from Russia, Germany, Holland, Morocco, and occasionally from Bombay, the last being oval and nearly twice as large as European Coriander, and pointed at the ends. A little is also cultivated in Essex. The English is rather pale and larger than the Dutch kind, which is small. The Russian is smaller than the other kinds. For distillation the fruits need to be previously crushed so as to expose the vittæ, which are found only on the inner surface of the mericarps. The powder of the fruits as seen under the microscope is characterised by the obliquely arranged linear cells of the endosperm, longer than those of Fennel, the sharply defined six-sided cells of the inner coat of the vittæ, and by the thick pitted walls of the large sclerenchymatous cells of the mesocarp. *J. Moeller, Leitfaden Mikr. Pharm. Übungen*, 1901, p. 182.

Tests—Coriander Fruits yield from 5 to 6 p.c. of ash. Five samples examined in the author's laboratory showed 4.69, 5.28, 5.74, 5.15 and 5.8 p.c., four samples of powdered Coriander gave 5.64, 5.7, 7.09, and 7.79. An ash limit of 6.0 p.c. for the fruits has been suggested.

Preparation

OLEUM CORIANDRI OIL OF CORIANDER

A colourless or pale yellow oily liquid, possessing a strong distinctive aromatic odour and taste.

It should be kept in well-closed glass bottles of a dark amber tint, and protected as far as possible from contact with air and light, and in a cool atmosphere.

Consists to the extent of 90 p.c. of dextrorotatory Linalool, $C_{10}H_{18}O$, sp. gr. 0.868, boiling point 194° to $198^{\circ}C$ (381° to $388^{\circ}F$).

Yield of Volatile oil from Coriandri Fructus—Moravian, Thuringian and Russian fruits yield, according to Gildemeister and Hoffmann, from 0.8 to 1.0 p.c. of oil, French, 0.4 p.c., Dutch, 0.6 p.c., Italian, 0.5 p.c., Moroccan, 0.2 to 0.3 p.c., whilst the East Indian fruit yields only 0.15 to 0.2 p.c.

Solubility—2 in 1 of Alcohol (90 p.c.), 1 in 75 of Alcohol (60 p.c.).

Used to render medicines more palatable, and prevent griping.

Dose— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Foreign Pharmacopœias—Official in U.S. Not in the others.

Tests—Coriander Oil has a specific gravity of from 0.870 to 0.885, is optically dextrogyrate, the rotation amounting to $+8^{\circ}$ to $+14^{\circ}$ in a 100 mm. tube. It yields a clear solution with three times its volume of Alcohol (70 p.c.).

The more generally occurring sophistications are sweet Orange Oil or Turpentine Oil, which are recognised by their influence on the specific gravity and the optical rotation, the solubility test affording additional evidence of the absence of Oil of Turpentine and added terpenes.

Not Official

COTO.

A Bark from Bolivia, origin unknown

Medicinal Properties—Aromatic stimulant and intestinal astringent Has been used in chronic diarrhoea

A precipitate may form when the Tincture or Fluid Extract is diluted with Water, but it diffuses readily, on being shaken, without the aid of Mucilage

It contains a bitter principle, **Cotoin**, sparingly soluble in Water, soluble in Alcohol, dose, $\frac{1}{2}$ to 2 grains = 0.03 to 0.13 gramme

Cotoin is recommended as checking the night sweats of phthisis —*L* '96, 1 255

Paracotoin is obtained from an allied bark, which has similar properties sparingly soluble in Water, soluble in Alcohol

Dose —2 to 3 grains = 0.13 to 0.2 gramme

Descriptive Notes—Coto bark as originally introduced into commerce is at present not obtainable, its place being taken by Paracoto bark. Both these barks come from Bolivia, the former from the banks of the Magdalena, and the latter from those of the Mapiri River, and are probably derived from trees belonging to the natural order Lauraceæ, so far as can be judged from their structure. Both are hard, heavy, reddish-brown barks occurring in quilled pieces about 6 to 8 inches (15 to 20 cm) long or more, 2 to 2½ inches (5 to 6 cm) broad, and about ½ inch (12.5 mm) thick, with a fracture granular outwardly and coarsely fibrous toward the inner surface. The chief difference between the two is that the inner surface in the true Coto bark is nearly smooth and the bark is thinner, that of Paracoto often being 15 mm or more thick and having the inner surface rough with projecting longitudinal ridges formed of sclerenchymatous fibres. Both have a pungent aromatic taste and exhibit a minute crystalline efflorescence on the broken surface. The taste of true Coto is rather more peppery than that of Paracoto. The distinctive microscopical characters are the large elongated sclerenchymatous cells, some of which contain granular matter, large parenchymatous cells containing yellow resin, and small simple starch granules. True Coto bark also contains granular reddish-brown oily bodies in some of the parenchymatous cells. The two barks may also be distinguished by the use of concentrated Nitric Acid, which turns Paracoto powder yellowish, changing gradually into a green tint, whilst that of Coto becomes deep red. This reaction is due to the Paracotoin and Cotoin.

TINCTURA COTO—1 of bruised Coto Bark macerated with Alcohol (90 p.c.) to make 10 —*B.P.C. Formulary* 1894 Incorporated in the *B.P.C.*

Dose —10 to 30 minims = 0.6 to 1.8 c.c.

Fluid extract (1 in 1), dose, 5 to 20 minims

Fortoin (Névalenc-Dicotoin) —Yellow, crystalline needles, or a light yellow powder. Insoluble in Water, soluble in Chloroform and Acetone. Decomposed by Alkali. Introduced as an intestinal antiseptic. Has been found useful in intestinal catarrh —*P.J.* '99, 11 168, '01, 1 702

Dose —1 to 5 grains = 0.06 to 0.32 gramme

CREOSOTUM.

CREOSOTE

Fr., CRÉOSÔTE OFFICINALE, GER., KREOSOT, ITAL., CREOSOTO, SPAN., CREOSOTA

A colourless or more generally a pale yellowish, highly refractive, oily liquid, possessing a strong phenolic odour and a burning, caustic taste. It is a mixture of Guaiacol, Creosol, and other Phenols obtained in the distillation of Wood Tar.

The *BP* states that Creosote is 'obtained in the distillation of Wood Tar', the *USP*, 'obtained during the distillation of Wood Tar, preferably that derived from the Birch'

It preserves animal substances from decay, from which property its name is derived. It is to the presence of this substance that the process of smoking hams owes its efficacy.

The two chief constituents of Creosote are **Guaiacol** and **Creosol**, the first of which predominates in some specimens, and the second in others. Beechwood Creosote contains most Guaiacol, formerly it was stated to contain more than 60 p c, but when the demand for Guaiacol and its salts arose, the proportion in commercial Creosote dropped to 20 p c. It can now be obtained containing 50 p c.

Guaiacol is soluble 1 in 80 of Water, and mixes with Glycerin in all proportions. Creosol is soluble 1 in 150 of Water, and will not form a clear mixture with Glycerin in any proportion.

Solubility—Beechwood Creosote is soluble about 1 in 110 of Water and mixes in all proportions with Alcohol (90 p c), Absolute Alcohol, Ether sp gr 0.735 and 0.720, Glacial Acetic Acid, Chloroform, Benzol, and Petroleum Spirit, it also mixes with Glycerin in all proportions up to nearly 3 of Glycerin to 1 of Creosote, but on the further addition of Glycerin the mixture is turbid.

'English Creosote' differs from Beechwood Creosote in that it is not nearly so soluble in Water, and does not mix readily with Glycerin. It dissolves about 1 in 350 of Water, and forms a turbid mixture with an equal volume of Glycerin.

Medicinal Properties—Disinfectant and antiseptic. It resembles Carbolic Acid in action, but it is less poisonous. Given internally in gastric fermentation, in putrefactive diarrhoea, and with considerable success in phthisis with abundant fetid sputum (*see below*), for arresting nausea in hysteria, for obstinate sea-sickness, and the vomiting of pregnancy and phthisis. A lotion (8 minims to 1 oz) and the ointment are used for eruptions of a scaly character, for venereal ulcers, and in parasitic skin diseases, it relieves the itching in eczema, toothache, when depending on caries, is relieved by its application. As an inhalation in fetid bronchitis, phthisis, and pulmonary gangrene.

Employed by internal administration with considerable success in phthisis, commencing with 5 minims in 2 fl drms of Cod liver Oil three times daily after meals and gradually increasing till at the end of three or four weeks 30 to 60 minims or even 80 minims are being taken three times daily. It is said to have no tendency to bad effects even in such large doses. Should a patient be unable to take Cod liver Oil, the Creosote may then be prescribed in spirituous solution. If the best Beechwood Creosote be used and due care exercised in increasing the dose gradually, it will be found to produce good results without unpleasantness or risk.—*B M J* '98, i 144, 299, 1388.

One drop of Creosote at bedtime every night for juvenile incontinence of urine.—*B M J* '87, i 809. In diabetes 4 drops daily increased to 10 drops.—*L* '89, i 702. Intratracheal injection of Creosoted Oil (1 in 20) to aid the expulsion of false membrane after tracheotomy.—*B M J* '98, i 1381.

Successful in cases of tuberculosis in children by pills and drops.—*T G* '93, 766.

Hypodermic injection of Creosote and Guaiacol dissolved in sterilised Almond Oil, 1 in 5 or 1 in 15.—*L* '96, ii 371, *B M J* '95, ii 1488. Small doses in gastric affections.—*L* '97, ii 404. In habitual constipation.—*L* '97, ii 932. Enemata containing 8 minims of Creosote in 4 oz of Cod liver Oil in pleuro peritoneal tuberculosis in children.—*L* '97, i 159. In malarial intermittent fever 15 minims

rubbed into the axilla and covered up with Cotton-Wool produced free perspiration and lowered the temperature—*B M J* '96, i 18, '97, i 1832, *I M G* '96, 11, *T G* '96, 325

Subcutaneous injection the best means of administering large quantities—*B M J* '01, ii 219

Creosote, Guaiacol, and their congeners are stated (*L* '04, ii 1827) to be much less used now in the treatment of pulmonary phthisis than a few years ago. An interesting item on Creosote is that each Japanese soldier is expected to carry and take Creosote pills as a prophylactic against dysentery—*B M J* '04, ii 1327

It is of distinct value in the antiseptic treatment of pulmonary tuberculosis (*Edin Med Jour* '05, 463). It often relieves gastric catarrh and stimulates the appetite. It should be given immediately after or before food in 2 or 3 minims doses, beginning with two or three times a day and gradually increasing to three times that amount, in capsules, or dissolved in Cod Liver Oil.

Twenty minims of a mixture of equal parts of a 20 p c Alcoholic solution of Creosote and Spirits of Chloroform, used for an hour or so on the sponge of an inhaler, relieves the troublesome cough of pulmonary phthisis—*Edin Med Jour* '05, 465

Dose—1 to 5 minims = 0.06 to 0.3 c c

Ph Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes

Prescribing Notes.—Given in capsules or in pills made with Soap and Liquorice Powder (see p 454). When given as a draught or mixture it is best emulsified with Mucilage of Gum Acacia and given in Milk, or dissolved in Almond Oil, see 'Guttæ Creosoti' and 'Mistura Creosoti' (Squire). For hypodermic injection, alone or dissolved in Almond Oil. When mixed with Magnesia it forms a tasteless compound insoluble in Water. Orange, Juniper, and Fluid Extract of Liquorice have been used as flavouring agents.

Incompatibles—When prescribed in pills with Silver Oxide it explodes, unless previously diluted with some inert powder.

Official Preparations—Mistura Creosoti, Unguentum Creosoti

Not Official.—Aqua Creosoti, Elixir Créosoté, Guttæ Creosoti, Mistura Pilula Creosoti, Solutio Creosoti Composita, Vapor Creosoti, creosoti, Parogenum Creosoti, Vin Créosoté, Creosoti Carbonas, Creosoti Oleas, Creosoti Phosphas, Creosoti Tannas, Creosoti Valerianas, Salocreol, Taphosote, Phosphotal, and Pneumin. The preparations of Guaiacol will be found under that name.

Foreign Preparation.—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung, Ita, , , , Port, Russ, Span, Swed, Swiss and U S

Tests—Creosote has a specific gravity of 1.080 to 1.086, and should not be below 1.080, the *BP* states not below 1.079. The *USP* states not below 1.078 at 25°C (77°F), the *PG* not below 1.08. Its boiling point is between 200° to 220°C (392° to 428°F), between which temperature the greater portion of it distils. It is either optically inactive or but slightly dextrogyrate. It is neutral or only feebly acid towards Litmus paper. A 1 in 100 solution in Alcohol (90 p c) or a 1 in 200 aqueous solution yields with Ferric Chloride Test-solution a green coloration rapidly changing to reddish-brown. Mixed with 10 times its volume of a 1 in 5 Solution of Potassium Hydroxide in Absolute Alcohol it forms a solid crystalline mass.

The more generally occurring impurities are Phenol, Coal Tar Creosote, neutral oils, Cærolignol, and higher boilingpoint constituents of Wood Tar. The *BP* is not official for Phenol and less volatile constituents. The *Ammon* is not officially adopted.

for its distinction from Phenol, it being required to suffer no material diminution in volume when shaken with 5 times its volume of Ammonia Solution. The absence of less volatile liquids is officially ensured by the absence of a translucent stain when dropped on to filtering paper and exposed to a temperature of 100°C (212°F). The Ammonia test has been stated (*CD* '00, 11 156, *PJ* '00, 11 150) not to be of special value, inasmuch as the purest Creosote shows the greatest diminution of volume. The author has found the best differentiating test between Creosote and Phenols to be the insolubility of the former in diluted Glycerin, three measures of Glycerin (sp gr 1.260) is diluted with 1 measure of Water and 1 volume of the Creosote sample is shaken with 3 volumes of the diluted Glycerin, after complete separation, the volume of the Creosote layer is read off, the diminution roughly indicating the amount of soluble impurity. If the Glycerin layer be separated and diluted with Water, the Coal Tar acids may be extracted by agitation with Chloroform, thus permitting their further examination. The *USP* mixes equal volumes of the Creosote and 95 p.c. Glycerin Solution, stating that a clear mixture will result, from which, on the addition of one-fourth volume of Water, a layer of Creosote equal to, or greater than, the volume originally employed will separate. The *USP* and the *PG* use Sodium Hydroxide Solution as a test for the presence of neutral oils, the *USP* mixing the Creosote with not less than 5 nor more than 9 times its volume of Normal Volumetric Sodium Hydroxide Solution, the *PG* mixing it with $2\frac{1}{2}$ times the volume of Sodium Hydroxide Solution (15 p.c.), in each case a clear liquid is required to result, which remains clear on dilution with 50 c.c. of Water. Coal Tar Creosote may be detected by the solubility of the Creosote in hot Water, its subsequent behaviour on cooling, and the behaviour of the filtrate with Bromine Water, the Collodion test, and the Alcoholic Potassium Hydroxide test. The Ferric Chloride test is stated to give an indication, but its usefulness is by no means fully conceded. The behaviour of the sample with Petroleum Ether and freshly prepared Barium Hydroxide Solution forms a useful means of readily detecting the presence of Cærolignol and some other high boiling point constituents of Wood Tar. The test is described in the small type below.

Fractionation—When distilled most of it comes over between 392°F (200°C) and 428°F (220°C), *USP* and *PG*. The *BP* states that it distils between these temperatures. When cooled to -20°C (-4°F) it becomes gelatinous but does not solidify, *USP* and *PG*.

Bromine—The saturated aqueous solution of Creosote separated from the oily globules yields a reddish-brown precipitate with T.S. of Bromine, *PG* and *USP*.

Collodion—If 1 volume of Creosote be shaken with 1 volume of Collodion no gelatinous mass should be formed, *BP* and *PG*, no permanent coagulum should form when equal volumes of the liquid are stirred together, *USP*.

Potassium Hydroxide—If 1 c.c. of Creosote be mixed with 10 c.c. of a solution (1-5) of Potassium Hydroxide in Absolute Alcohol, a solid crystalline mass will form, *USP* and *PG*.

Benzin and Barium Hydroxide—If 1 c c of Creosote be shaken with 2 c c of Petroleum Benzin and 2 c c of Baryta Water, the Petroleum Benzin solution should not assume a blue colour or be muddy, and the aqueous liquid should not be coloured red, *P G*

If 1 c c of Creosote be cautiously and gently shaken with 2 c c of Petroleum Benzin and 2 c c of a freshly prepared Barium Hydroxide T S until of uniform consistence, on complete separation three distinct layers are visible, the middle one of which contains the Creosote unaltered in appearance, while the Petroleum Benzin should not be blue or muddy, and the aqueous layer should not have acquired a red tint, indicating the absence of Cærulignol and some other high boiling constituents of Wood Tar, *U S P*

Preparations

MISTURA CREOSOTI. CREOSOTE MIXTURE

Shake 16 minims of Creosote with 14 fl oz of Distilled Water, add 1 fl oz of Syrup and 16 minims of Spirit of Juniper, and Distilled Water, *q s* to yield 16 fl oz (1 in 480)

It was pointed out in the *Companion* that Glacial Acetic Acid was quite unnecessary, and it is now omitted

Dose.— $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

UNGUENTUM CREOSOTI.—CREOSOTE OINTMENT

Creosote (by weight), 1, Hard Paraffin, 4, Soft Paraffin, white, 5; add the Creosote to the melted Paraffins, and stir until cold (1 in 10)

Now made with Hard and Soft Paraffins in place of Simple Ointment

Not Official

AQUA CREOSOTI—Creosote, 10, Distilled Water, 990 Agitate the Creosote vigorously with the Distilled Water, and filter through a well-wetted filter Creosote Water should be freshly prepared when dispensed — *U S P*

This has been incorporated in the *B P C* under the title **Liquor Creosoti** with *syn* Aqua Creosoti

ELIXIR CRÉOSOTÉ—Creosote, 1, Rum, 66, mix and filter

Spiritus Creosoti—Creosote, 1, Alcohol (90 p c), 40 Dose — 1 dr m — *Martindale*

This has been incorporated in the *B P C*

GUTTÆ CREOSOTI (*Squire*)—Creosote, 16 minims, Mucilage of Gum Acacia, 60 minims, Syrup of Orange, 1 fl oz, Water, to 2 fl oz, mix the Creosote with the Mucilage and add the other ingredients One or two teaspoonfuls for a dose in an oz of Milk

MISTURA CREOSOTI (*Squire*)—Creosote, 16 minims, Almond Oil, $\frac{1}{2}$ fl oz, Syrup of Orange, 1 fl oz, Powdered Gum Acacia, $1\frac{1}{2}$ dr m, Water, to 8 fl oz Dissolve the Creosote in the Oil, mix it with the Powdered Gum Acacia in a mortar, add all at once 3 fl dr m of Water, and triturate until an emulsion is formed, then add the remainder of the Water and the Syrup of Orange

Dose.— $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

PILULA CREOSOTI—Creosote, 12 minims, Curd Soap, in powder, 6 grains, Liquorice, in powder, 30 grains, mix, and divide into 12 pills

This has been incorporated by the *B P C*

Foreign Pharmacopœias—Official in Austr, Creosote 5, Yellow Wax 2 5 Liquorice 6, Extract of Liquorice 6, Mucilage of Acacia *q s*, Belg, Creosote 10, Water 2, Liquorice Root *q s*, Ger, and Jap, Creosote 10, Liquorice 19, Glycerin 1, Swiss, Creosote 5, Liquorice 9 5, Glycerin 0 5 Fr, Creosote with

Medicinal Soap *g s* Each pill contains 10 centigrammes of Creosote, 0.1 gramme ($1\frac{1}{2}$ grains)

SOLUTIO CREOSOTI COMPOSITA—Creosote, 1, Spirit of Menthol (20 p c), 1, and Spirit of Chloroform, 1—*Brompton* Useful in an oro-nasal inhaler

VAPOR CREOSOTI (*B P* 1885)—Creosote, 12 minims, Boiling Water, 8 fl oz Mix the Creosote and Water in a suitable apparatus, for inhalation

This has been incorporated in the *B P C*

Creosote, 80 minims, French Chalk, 30 grains, Water, to 1 oz A teaspoonful in 20 oz of Water at 140° F for each inhalation—*Throat*

VASOLIMENTUM CREOSOTI—Creosote, 5, Liquid Vasoliment, 95—*Hager*

Parogenum Creosoti—Creosote, 5, Parogen, 95—*B P C*

VIN CRÉOSOTÉ—Creosote 1, Alcohol (90 p c) 9, Simple Syrup 10, Malaga Wine 80—*Fr*

CREOSOTI CARBONAS (Creosotal)—A viscid, amber coloured liquid, nearly odourless and tasteless, insoluble in Water Not only useful in chronic diseases of the lung, but in acute diseases of the respiratory organs It is stated to contain 90 p c of Creosote, and to be free from the irritating effects of that substance—*B M J E* '96, i 15, *L* '97, ii 1472

One teaspoonful doses for adults, smaller doses for children—*L* '98, i 222, this dose has been criticised, and 5 drops three times daily recommended—*L* '98, i 960

Is preferable to the Phosphate, though both are better than pure Creosote—*B M J* '01, ii 219

Teaspoonful doses morning and night, taken in a cup of hot sugared Milk in the treatment of acute broncho pulmonary affections The dosage for children is proportionately smaller Has remarkable power of reducing temperature in bronchitis and pneumonia, and is beneficial even in advanced pneumonia Administration best stopped gradually—*L* '99, ii 710, *B M J E* '02, i 4

Foreign Pharmacopœias—Official in Austr, Belg, Jap and Swiss

CREOSOTI OLEAS (Oleocreosote)—A light yellow, oily liquid, having a faint odour and taste of Creosote Insoluble in Water, soluble in Absolute Alcohol and in Ether

Dose—15 to 30 grains = 1 to 2 grammes

CREOSOTI PHOSPHAS (Phosphote)—A dense, oily substance, insoluble in Water

Dose—5 to 15 grains in capsules = 0.32 to 1 gramme

CREOSOTI TANNAS (Tannosal)—A brown, hygroscopic powder, soluble in Water, in Alcohol (90 p c) and in Glycerin

Dose—5 to 15 grains = 0.32 to 1 gramme

CREOSOTI VALERIANAS (Eosote)—A yellow, oily liquid, distilling at 240° C (464° F), insoluble in Water, soluble in Alcohol (90 p c) and in Ether Has been recommended as a substitute for Creosote on account of its freedom from corrosive and toxic properties Commencing dose 3 grains, increasing to 6 or 9 grains three times a day, given in capsules—*B M J E* '96, ii 59

SALOCREOL (Creosote Salicylic Ester)—A brown, oily, neutral liquid, insoluble in Water, readily soluble in Alcohol (90 p c), in Ether and in Chloroform It has been used in the treatment of rheumatic swelling of the joints—*B M J E* '08, ii 52

Dose—6 to 20 grammes rubbed into the skin

Taphosote, the Tannophosphoric Ester, and **Phosphotal**, the Phosphite, are combinations of Phosphoric Acid and Creosote

Pneumin, a compound of Creosote and Formaldehyde, is a yellow, tasteless, odourless powder Insoluble in Water Stated to have a beneficial effect in tuberculosis

Dose— $7\frac{1}{2}$ to 80 grains = 0.5 to 2 grammes

CRETA PRÆPARATA.

PREPARED CHALK

A purified native Calcium Carbonate, most of the impurities having been removed by elutriation

Solubility—Insoluble in Water, readily dissolved by weak acids

Medicinal Properties—It is astringent and antacid Combined with other astringents and aromatics, it is used in infantile diarrhoea and in diarrhoea accompanied with acidity One of the best antidotes for Oxalic Acid, the mineral acids, and Zinc Chloride Used as a dusting powder in burns, ulcers, and moist eczema, it is protective and desiccant

Dose—10 to 60 grains = 0.65 to 4 grammes

Prescribing Notes—Generally given in the form of *Mistura Cretæ* with astringent Tinctures and Opium

The Pulvis Cretæ Aromaticus is useful for administration to children, either in powder or in mixture with Mucilage

Incompatibles—All Acids and Sulphates

Official Preparations—*Mistura Cretæ*, *Pulvis Cretæ Aromaticus* and *Pulvis Cretæ Aromaticus cum Opio* Contained in *Hydrargyrum cum Cretâ*

Not Official—Cholera Mixture, *Pulvis Cretæ Compositus* and *Unguentum Cretæ*

Foreign Pharmacopœias—Official in Hung, Jap, Port, Span and U S Not in the others

Tests—Prepared Chalk is dissolved readily by dilute acids, effervescence occurring, with the evolution of a colourless and odourless gas, which affords a white precipitate when passed into Lime Water A solution prepared by dissolving a portion of the sample in just sufficient Hydrochloric or Nitric Acid to effect solution, boiled and cooled, answers the tests distinctive of Calcium given under Precipitated Calcium Carbonate

The more generally occurring impurities are siliceous material, Iron, Aluminium, Phosphates, Sulphates and Barium Carbonate Siliceous material may be detected by remaining insoluble in Hydrochloric Acid, preferably after the acid solution to dryness and re-solution in diluted Hydrochloric Acid, and which should be relatively minute Iron, Aluminium, Magnesium, Phosphates and Sulphates may be examined for by the tests given under *Calci Carbonas Præcipitatus*, and should be present in but slight traces Barium Carbonate, if present may be detected by dissolving a portion of the sample in Diluted Acetic Acid and adding Potassium Chromate Solution, a yellow precipitate insoluble in Acetic Acid, soluble in diluted mineral acids, indicates the presence of Barium

Preparations

MISTURA CRETÆ. CHALK MIXTURE

Prepared Chalk, $\frac{1}{2}$ oz, Tragacanth, in powder 15 grs Refined Sugar, $\frac{1}{2}$ oz, Cinnamon Water, q s to make 8 fl oz (about 1 in 32)

Tragacanth is now used in place of Gum Acacia, and Sugar in place of Syrup

Dose — $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

Foreign Pharmacopœias — Official in Port, Carbonate of Lime 3, Gum Arabic 3, Syrup of Cinnamon 10, Water 84, U S, Prepared Chalk 6, Acacia 4, Sugar 10, Cinnamon Water 40, Water, to measure 100 Not in the others

PULVIS CRETÆ AROMATICUS AROMATIC POWDER OF CHALK

Prepared Chalk, 11, Cinnamon, 4, Nutmeg, 3, Cloves, $1\frac{1}{2}$, Cardamom Seeds, 1, Refined Sugar, 25, all in powder
(about 1 Chalk in $4\frac{1}{2}$)

Saffron is now omitted

Dose — 10 to 60 grains = 0 65 to 4 grammes

PULVIS CRETÆ AROMATICUS CUM OPIO AROMATIC POWDER OF CHALK WITH OPIUM

Aromatic Powder of Chalk, 39, Opium, in powder, 1
(1 Opium in 40)

Dose — 10 to 40 grains = 0 65 to 2 6 grammes

Not Official

CHOLERA MIXTURE — Aromatic Powder (*B P* '64), 3 dr̄m, Spirit of Sal Volatile, 3 fl dr̄m, Tincture of Catechu, 10 fl dr̄m, Compound Tincture of Cardamoms, 6 fl dr̄m, Tincture of Opium, 1 fl dr̄m, Chalk Mixture, to make 20 fl oz

Dose — 1 fl oz = 28 4 c c for an adult, $\frac{1}{2}$ fl oz = 14 2 c c for a child 12 years old, $\frac{1}{4}$ fl oz = 7 1 c c for seven years old, after each liquid motion

This mixture was proposed by the Board of Health during the prevalence of cholera, and is useful in cases of diarrhœa

A mixture much like this has been introduced into the *B P C* under the title **Mistura Cretæ Composita** with the *synonym* Board of Health Cholera Mixture as follows —

Compound Aromatic Powder, 2, Aromatic Spirit of Ammonia, 1 875, Tincture of Catechu, 6 25, Compound Tincture of Cardamoms, 3 75, Tincture of Opium, 0 625, Chalk Mixture, *q s* to produce 100

PULVIS CRETÆ COMPOSITUS — Prepared Chalk, 30, Acacia, in fine powder, 20, Sugar, in fine powder, 50 — *U S P*

This has been incorporated in the *B P C*

UNGUENTUM CRETÆ — Prepared Chalk, 1, Spermaceti Ointment, 4, mix

CROCUS.

SAFFRON

FR, SAFRAN, GER, SAFRAN, ITAL, ZAFFERANO, SPAN, AZAFRAN

The dried Stigmas and tops of the Styles of *Crocus sativus*

Imported from Spain, France, and Italy

The important ingredient of Saffron is *Crocine* or *Polycrocin*, a body of a glucosidal nature stated by Hilger and Schuler to be identical with Carotin. It also contains a small percentage from 0 75 to 1 0 p c of Volatile Oil and Picrocrocin, in addition to Wax, Gum, Albumen, saline matter, Water, and Lignin

Medicinal Properties — Useful for giving colour and flavour to preparations

tum Aloes Compositum and Tinctura Cinchonæ Composita

Not Official—Glycerinum Croci and Syrupus Croci

Foreign Plasmid Official in all except US , Dutch, Stigmata
Croci, Mex, Arabian, Arab, Acafrao, Dan, Norw and Swed, Stigma
Croci

Descriptive Notes—Saffron consists of the upper part of the trifid style, and stigmas of *Crocus sativus*, L. It is of an orange-red colour and is usually about one inch or more in length, the stigmatic portion being slightly dilated, nearly tubular above, slit on the under side, and toothed at the apex. In commerce it is frequently adulterated, no drug more so. The best and purest commercial variety is that from Valencia, the Saffron of Alicante, and particularly of Barcelona, being often adulterated. The stamens of the flower are sometimes present in more than accidental amount, and are even sometimes offered separately in the drug market as 'yellow saffron'. They are linear and arrow-shaped at the base, inserted on a short filament. The florets of *Calendula officinalis*, L., are seen to be flat and pale in colour, and the roots of *Carex* thread-like. Saffron normally retains about 12½ p.c. of moisture (12 p.c., P.G.), and should therefore be kept in a tin, or loss of weight may ensue. The fresh crop is usually obtainable in commerce in November and December.

The florets of *Canthamus tinctorius*, L. (nat. ord. *Compositae*), sold in small flat cakes, are sometimes offered as 'Cake Saffron'. The florets are saffron-coloured, tubular, and contain syngenesious anthers

Under the name of Cape Saffron the flowers of *Lyperna crocea*, Eckl (nat ord *Scrophulariaceae*), are at rare intervals offered in the drug market. It is a native of the Cape of Good Hope.

Tests—Saffron yields an intensely yellow solution when treated with Water, and when rubbed on the wet finger leaves an orange-yellow stain. The *PG* states that 100,000 parts of Water shaken with 1 part of Saffron assume a pure and distinct yellow colour. When brought into contact with a drop of Sulphuric Acid it yields a deep indigo-blue coloration.

The more generally occurring adulterations of Saffron are excess of moisture, mineral matter, e.g., Barium Sulphate, Sand, etc., fixed oils, stamens artificially dyed so as to resemble stigmata, &c. Nitrates, due to the presence of artificial colouring matter derived from Nitrophenols or Nitroresols, principally the latter. The limit of moisture is fixed by the *BP* at not more than 12.5 p.c., by the *PG* at not more than 12 p.c. Mineral matter may be detected by the residue left on ignition, or by a deposit of white or coloured powder settling out when the sample is floated on the surface of some warm water. The finest Saffron yields, when ignited with free access of air, from 4.4 to 5.5 p.c. of ash. The *BP* limit is about 7.0 p.c., that of the *PG* calculated on dried sample 6.5 p.c., which is equivalent to 7.4 p.c. on the undried Saffron. The ash should be examined for Barium.

The ash is considered (C D '02, 1 118) to be of little use to the buyer of saffron unless he has experience as well to help him, a poor, thin, semi-wild

variety of Saffron may give an excellent ash, possibly not over 4 p c, whilst a fine bold quality which has been dressed to a very moderate extent will be outclassed by yielding over the *B P* maximum

Fixed oils may be detected by the greasy spot produced when the sample is pressed between folds of white bibulous paper, artificially dyed products by the Sulphuric Acid test after extraction with Petroleum Ether, which extracts the colour derived from Coal Tar products but not that of genuine Saffron Safflower yields an Infusion which is coloured greyish green by Ammonia Solution and a bright red by Nitric Acid Colouring matters derived from Nitrophenols or Nitro-cresols may be detected by deflagration occurring during incineration, or by dissolving the ash in Water and applying the Ferrous Sulphate test for Nitrates The colouring powers of different specimens of Saffron may be judged by comparison with a standardised Potassium Bichromate Solution

Preparation

TINCTURA CROCI TINCTURE OF SAFFRON

1 of Saffron, macerated with 20 of Alcohol (60 p c) (1 in 20)

Dose —5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias —Official in Spain, 1 in 5, Belg, Jap and Swiss, 1 in 10, all by weight Not in the others

Tests —Tincture of Saffron has a specific gravity of 0.920 to 0.925, contains from 2.0 to 3.0 p c w/v of total solids and from 57 to 58 p c w/v of Absolute Alcohol A few drops of the Tincture evaporated to dryness in a white porcelain dish on a water-bath leave a residue which when cooled yields an indigo-blue coloration with a drop of concentrated Sulphuric Acid

Not Official

GLYCERINUM CROCI (*Squire*) —Saffron, 1, Glycerin, 20, Alcohol (60 p c), 20, mix the Glycerin and the Alcohol, and digest in it the Saffron for an hour at a gentle heat, and filter This is introduced as a substitute for Syrupus Croci, which deposits and loses its colour

This has been incorporated in the *B P C*

SYRUPUS CROCI (*Squire*) —Glycerin of Saffron, 1, Syrup, 7

This has been incorporated in the *B P C*

CROTONIS OLEUM.

CROTON OIL

FR, HUILE DE CROTON, GER, KROTONÖL, ITAL, OLIO DI CROTONTIGLIO,
SPAN, ACEITE DE CROTON TIGLIO

A yellow, brownish yellow, or reddish-brown, somewhat viscid, slightly fluorescent, oily liquid, possessing an unpleasant odour and an acid and burning taste It is the Oil expressed from the seeds of *Croton Tiglium*, L

It consists chiefly of the Glycerides of Stearic, Palmitic, Myristic, Lauric, and Oleic Acids It also contains the Glycerin Esters of Formic, Acetic, Isobutyric,

and Isovaleric Acids, together with Tiglic Acid, Crotonoleic Acid, a toxic anisomeric Croton, and Croton-resin, the latter lactone possessing powerful vesicant properties.

A native of Hindostan, Ceylon, and the Moluccas

100 of seed yield about 50 of Oil

Solubility—Soluble in Ether, Oil of Turpentine, and Olive Oil, partially soluble in Alcohol (90 p c)

B P 1898 still retains the sentence, 'entirely soluble in Absolute Alcohol,' although it has been repeatedly pointed out that this is not strictly the case. An oil recently expressed will dissolve the Absolute Alcohol up to equal parts, but when more than one volume of Alcohol is added to one of Oil the mixture becomes turbid, and with two volumes of Alcohol the mixture separates into two layers on standing. With a sample of oil two or three years old rather more Alcohol can be added without the mixture becoming turbid, but it is only a question of degree.

The solubility of Croton Oil in Absolute Alcohol appears to depend in great measure on the age of the Oil, and the greater or less freshness of the seeds from which it is expressed, as oxidised or resmified Oil dissolves more readily.

The solubility of the Oil as a whole depends upon the proportion of free Acid, which is very soluble in Alcohol, and also carries the difficultly soluble neutral Glyceride into solution along with it.

Croton Oil can be separated by Alcohol into two parts. The non-vesicating portion insoluble in Alcohol possesses the full purgative properties of the Oil in a solid form, the alcohol-soluble or vesicating portion had no purgative same doses, but caused irritation and nausea.

Medicinal Properties—A powerful drastic cathartic, acting with great rapidity. Given in cases of obstinate constipation, in dropsy, in apoplexy, in maniacal and unconscious patients, and in eclampsia, its small dose being an advantage. Applied externally as a powerful counter-irritant in rheumatism, gout, neuralgia, and in acute laryngeal and pulmonary diseases in the form of liniment. Its external application is painful, and is often followed by an inflammatory eruption which becomes pustular, and leaves unsightly scars. It is therefore not often used externally, unless well diluted.

Croton Oil must be given with great care, and is inadmissible in feeble subjects, in organic obstruction, and in inflammatory states of the stomach and intestines.—*Mitchell Bruce*

It should never be given to children, to pregnant women, to those with hemorrhoids, nor to those suffering from peritonitis.—*Hale White*

5 minims to 1 fl oz of Olive Oil are used to promote the growth of hair

Dose— $\frac{1}{2}$ to 1 minim = 0.03 to 0.06 cc

Ph Ger maximum single dose, 0.05 gramme, maximum daily dose, 0.15 gramme

Prescribing Note—*I will with Soap and Liqueur Powder (see p 897), or in emulsion with C. Extract of Colocynth*

Official Preparation—Linimentum Crotonis

Not Official—Croton Oil Pencils and Collodium Tiglin

Antidotes—In case of an overdose an emetic should be at once administered, the stomach should be washed out with 1 oz to pint of Water, mucilaginous fluids and Opium or 1 to 2 gr of Morphia be given to check the pain and enteritis.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fl, Ger, Hung, Ital, Jap, Mex (Aceite de Croton Tiglio) Norw, Port, Russ, Span, Swed, Swiss and US (Oleum Tiglin). Swed has also an Oleum Crotonis Extractum.

Tests—Croton Oil has a specific gravity of 0.940 to 0.960 which limits are given by both the *BP* and the *PG*, the *USP* gives 0.935 to 0.950 at 25° C (77° F). It should be slightly dextrogyrate. It possesses an Acid value of 21 to 22, a Saponification value of 203 to 215, and an Iodine absorption of not less than 105.0 p.c. A sample of the fresh oil examined in the author's laboratory had an Acid value of 24.36, an Ester value of 176.4, a Saponification value of 200.76, a sample dated 1906 had an Acid value of 20.9, an Ester value of 179.2, a Saponification value of 200.1, and an Iodine value of 106.68, a sample dated 1900 had an Acid value of 48.72, an Ester value of 149.8, and a Saponification value of 198.52. The solidifying point of the fatty acids ranges from 16.5° to 16.8° C (61.7° to 62.24° F). The *BP* refers to the specific gravity, and states that the alcoholic solution should not redden blue Litmus paper, but makes no reference to the other physical and chemical characteristics of the oil.

Croton Oil may be detected in mixtures by shaking with an Alcoholic Potassium Hydroxide solution, separating the alcoholic layer, acidifying with dilute acid and removing the spirit by distillation. The residue is shaken with Ether, the ethereal solution separated, the Ether distilled and the residue tested on the skin. A characteristic pustular eruption should be produced if Croton Oil be present.

The more generally occurring impurities are 'other non-drying oils'. These are detected by vigorously shaking a measured quantity of 2 c.c. of the oil with a mixture of 1 c.c. of fuming Nitric Acid and 1 c.c. of Water, after standing for one or two days the mixture should neither partially nor completely solidify. The test is common to the *BP*, *USP* and *PG*.

Saponification—The *USP* requires that Croton Oil should show a Saponification value of from 203 to 215 when saponified by Alcoholic T.S. of Potassium Hydroxide.

Iodine Absorption—If 0.8 gramme of Croton Oil be dissolved in 10 c.c. of Chloroform in a 250 c.c. bottle or flask, and 25 c.c. of a mixture of equal volumes of Alcoholic Iodine T.S. and Alcoholic Mercuric Chloride T.S. added, and if after standing for 4 hours protected from light 20 c.c. of Potassium Iodide T.S. be added and the mixture diluted with 50 c.c. of Water, on titrating the excess of Iodine with Tenth normal V.S. of Sodium Thiosulphate, an Iodine value of not less than 103 nor more than 109 should be obtained, *USP*.

Preparation

LINIMENTUM CROTONIS LINIMENT OF CROTON OIL

Croton Oil, 1, Oil of Cajuput, $3\frac{1}{2}$, Alcohol (90 p.c.), $3\frac{1}{2}$ (1 in 8)

Brompton and *St. Mary's* have a diluted liniment made with equal parts of the Official Preparation and Liniment of Soap.

Not Official

CROTON OIL PENCILS—Croton Oil, 2, Cacao Butter, 1, White Bees wax, 1, melt together the last two in a water-bath, add the Oil, and when nearly cold pour into moulds.

COLLODIUM TIGLII—Croton Oil, 1, Flexible Collodion, 9—*USNF*

CUBEÆ FRUCTUS.

CUBEBS

FR, CUBÈBE, GER, KUBEEN, ITAL, PEPE CUBESE, SPAN, CUBEBA.

The dried, full-grown, unripe Fruits of *Piper Cubeba*

Medicinal Properties—Stimulant and antiseptic diuretic, expectorant Acts specially on the genito-urinary mucous membrane Given in all stages of gonorrhœa, gleet, cystitis, pyelitis, and sometimes in chronic bronchitis Frequently combined with Copaiba

Dose—30 to 60 grains = 2 to 4 grammes

Prescribing Notes—The Powder is given in the above moistened wafer paper, or in smaller doses in cachets In mixture well rubbed down with Mucilage A popular form of administration is the paste, made with an equal quantity of Copaiba, which may be taken in wafer paper It is also made into a paste with Glycerin and various Syrups For throat affections, Lozenges, Compressed Tablets, and Cigarettes are made It is also given in the form of Vapour

The Oil is given in Capsules or suspended in Water with Mucilage For Inhalation the Oil may be used with or without the vapour of Water

Official Preparations—Oleum Cubebæ and Tinctura Cubebæ

Not Official—Extrait de Cubèbe, Fluidextractum Cubebæ, Oleo-resina Cubebæ, Trochiscus Cubebæ, Vapor Cubebæ and Vapor Cubebæ cum Limone

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Descriptive Notes—Cubebs vary much in quality, and when scarce and dear are often adulterated, or other species are substituted for them The official kind should be about $\frac{1}{8}$ inch (4 mm) in diameter and vary from nearly black to greyish-brown in colour, the pericarp being wrinkled, and furnished with a slender rounded pedicel about $1\frac{1}{2}$ times as long as the fruit and continuous with it (4 to 10 mm, P G) The pericarp contains a single seed attached to the base of the ovary The taste is warm and aromatic and slightly bitter These characters are, however, not sufficient to distinguish the genuine from the false fruits frequently offered in commerce The most reliable characters, adopted in the B P are (1) the crimson colour developed when a crushed Fruit is covered with a drop or two of Sulphuric Acid, since the spurious Cubebs hitherto met with do not contain Cubebin and Cubebic Acid, to which the reaction is due, (2) in the Powdered Fruit, the radially elongated cells of the inner surface of the pericarp Three varieties of the plant are cultivated in Java The Fruit of two of them give the crimson reaction with Sulphuric Acid, but the Fruit of one is longer than that of the other, the third kind has an odour recalling that of Nutmegs, it does not give the crimson reaction, and has caused symptoms of poisoning when administered, see P J (3), xxv, pp 314, 757, 797

When Cubebs are scarce they are adulterated with Fruits similar in form, such as *Rhamnus* species, which is a 3- to 4-celled Fruit, *Bredelia montana*, Willd, and *Litsea citrata*, Bl, which have a large exalbuminous embryo, whilst that of *Piper Coccia* is minute and embedded in the apex of a large oily and barbaous perisperm

Other species of Piper Fruits are sometimes substituted for Cubebs, *e.g.*, *Piper ribesoides*, Wall, *P. crassipes*, Korth, and *P. Lowong*, Bl, but these are either larger than Cubebs or are different in flavour, and do not give the crimson reaction with Sulphuric Acid

The quality of genuine Cubebs depends upon freedom from stalks, and from immature hollow Fruits, which are concave at the base, since the stalks or rachis contain less Oil, and the Seed contains more, than the pericarp. Usually the stalks are sold separately, and employed for the distillation of Oil of Cubebs

Tests—Cubebs when crushed and tested with Sulphuric Acid impart a crimson colour to the Acid. No limit of ash is given in the *BP*, it should not exceed 7.0 p.c.

An Oleo-resin extractable by Ether and by Alcohol (90 p.c.) is present in the Fruits, to the extent of from 17 to 25 p.c. A limit of 22.0 p.c. has been suggested (*CD '02*, 11 826) as a standard for inclusion in the official monograph, dry chemically pure Ether being suggested as a solvent. Ether was employed as a solvent in the 1890 Edition of the *USP*, but was altered to Alcohol (94.9 p.c.) in the 8th Decennial Revision. A standard of not less than 17.0 p.c. has also been suggested

Preparations

OLEUM CUBEBAE OIL OF CUBEBS

A viscid, oily liquid, possessing a characteristic odour and a warm camphoraceous taste

BP describes it as colourless, pale green, or greenish yellow, Schimmel, as light green, or bluish green, it is colourless only when the last portions of the distillation, which are blue, have not been added to the product

It is distilled from Cubebs, the yield being from 10 to 18 p.c.

It consists almost entirely of Terpenes or Sesquiterpenes. It contains a laevogyrate Terpene, Pinene, Dipentene, a laevogyrate Sesquiterpene, Cadinene, and a laevogyrate Sesquiterpene Alcohol, Cubebs-camphor, which is found only in old Oil

Solubility—1 in 18 of Alcohol (90 p.c.), in all proportions of Absolute Alcohol

Dose—5 to 20 minims = 0.3 to 1.2 c.c.

Foreign Pharmacopœias—Official in *Port*, sp. gr. 0.929, *US*, sp. gr. 0.905 to 0.925 at 25° C (77° F). Not in the others

Tests—Oil of Cubebs has a specific gravity of from 0.915 to 0.930, the *USP* gives 0.905 to 0.925 at 25° C (77° F), the *BP* 0.910 to 0.930. It has an optical rotation of -25° to -40° in a 100 mm tube. The greater portion of the Oil distils between 250° and 280° C (482° and 536° F), about 10 p.c. passing over below 250° C (482° F). The Oil is stated to be soluble in from 1 to 3 volumes of Alcohol (90 p.c.), and to afford a solution which is neutral to Litmus paper. The solubility in Alcohol (90 p.c.) varies greatly according to the age of the sample, old Oils being apparently more soluble than new Oils. The Oil is not often adulterated. Turpentine Oil if present would be detected by the behaviour of the Oil on fractionation

TINCTURA CUBEBÆ. TINCTURE OF CUBEBS

4 of Cubebs, percolated with Alcohol (90 p c), to yield 20 (1 in 5)

Dose — $\frac{1}{2}$ to 1 fl drm = 18 to 36 c c

Foreign Pharmacopœias — Official in Mex, 1 in 5, by weight Not in the others

Tests — Tincture of Cubebs has a specific gravity of about 0.840, contains about 20 p c w/v of total solids and about 86.0 p c w/v of Absolute Alcohol

Not Official

EXTRAIT DE CUBÈBE — 1 of Cubebs, in No. 22 powder, percolated first with 2 of Ether, and subsequently with 2 of Alcohol (95 p c), evaporate the two liquids separately and mix the resulting extracts — *F*

FLUIDEXTRACTUM CUBEBÆ — Cubebs, in No. 40 powder, 100 grammes, percolated with Alcohol (95 p c by vol) until the Cubebs are exhausted, reserve the first 90 c c of percolate, and evaporate the remainder to a soft extract, dissolve this in the reserved portion, and add sufficient Alcohol to make 100 c c — *U S P*

This has been incorporated in the *B P C*

Dose — 5 to 30 minims = 0.3 to 1.8 c c

OLEO-RESINA CUBEBÆ *Syn* **EXTRACTUM CUBEBÆ** — Percolate Cubebs in coarse powder with Ether, slowly, until the liquor passes colourless. Let the Ether evaporate from the liquor, at first spontaneously and then over a water-bath, or recover it by distillation, and transfer the residue to a closed vessel, letting it stand until waxy or crystalline matter ceases to be deposited. Decant the Oleo-resin and preserve it in a well stoppered bottle.

This was official in *B P* 1885, and has been incorporated in the *B P C*

Dose — 5 to 30 minims = 0.3 to 1.8 c c

Foreign Pharmacopœias — Official in Fr, Ger, Hung, Jap, Mex (*Extracto Alcohólico de Cubebas*), Swiss and U S Not in the others

TROCHISCUS CUBEBÆ — Each lozenge contains about $\frac{1}{2}$ grain of Cubebs with Fruit Paste — *Throat*

This has been incorporated in the *B P C*

Official in U S, about $\frac{1}{2}$ grain of Oleo resin in each

VAPOR CUBEBÆ — Oil of Cubebs, 40 minims, Light Magnesium Carbonate, 20 grains, Water, to 1 fl oz Mix. A teaspoonful in a pint of Water at 140° F for each inhalation — *Throat*

This has been incorporated in the *B P C*

VAPOR CUBEBÆ CUM LIMONE — Oil of Cubebs, 30 minims, Oil of Lemons, 10 minims, Light Magnesium Carbonate, 20 grains, Water, to 1 oz — *Throat*

Not Official**CUCURBITÆ SEMINA PRÆPARATA**

MELON PUMPKIN SEEDS |

The prepared fresh ripe Seed of *Cucurbita maxima*, from cultivated plants are official in the *Ind* and *Col* Add for the Mediterranean Colonies

Dose — 3 to 4 oz = 85.2 to 113.6 grammes

Not Official
CUPRI SUBACETAS

Syn—ÆRUGO VERDIGRIS

Pale green powder, or partly crystalline masses

According to Von Hager two varieties are recognised commercially the blue or French Verdigris, consisting chiefly of monobasic Copper Acetate, $\text{Cu}(\text{C}_2\text{H}_3\text{O})_2 + \text{Cu}(\text{OH})_2 + 5\text{H}_2\text{O}$, and the green, or English, German, or Swedish variety, consisting chiefly of semi basic Copper Acetate $[\text{Cu}(\text{C}_2\text{H}_3\text{O}_2)_2] + \text{Cu}(\text{HO}) + 5\text{H}_2\text{O}$, as well as some bibasic Copper Acetate $\text{Cu}(\text{C}_2\text{H}_3\text{O}_2)_2 + 2\text{Cu}(\text{HO}) + \text{H}_2\text{O}$

Solubility—When treated with Water about 50 p c dissolves as Copper Acetate, leaving an insoluble Acetate insoluble in Alcohol (90 p c), soluble in diluted mineral acids and in Acetic Acid, also soluble in Ammonia

Medicinal Properties—Used as a stimulant to foul and indolent ulcers, also as an escharotic

Foreign Pharmacopœias—Official in Belg, Mex (Acetato de Cobre bibasico), Port (Verdete), and Span (Cardenillo) Not in the others

Tests—Copper Subacetate answers the tests for Copper appearing under Copper Sulphate When warmed with Sulphuric Acid and a little Alcohol (90 p c), the distinctive odour of Ethyl Acetate is evolved, when warmed with a minute amount of Arsenious Anhydride, the characteristic and highly poisonous odour of Cacodyl Oxide is evolved It should dissolve almost completely in Ammonia Solution

The more generally occurring impurities are Arsenic, metallic Copper, Aluminium, and Chalk Arsenic may be detected by Bettendorf's test, metallic Copper and Aluminium from the residue insoluble in Ammonia Solution, and Chalk by the effervescence on the addition of Hydrochloric Acid Copper Sulphate, if present, may be detected by the addition of Barium Chloride solution

LINIMENTUM ÆRUGINIS (*Ph Lond*)—Made by dissolving Verdigris 1, in Vinegar 7, adding Honey 14, and boiling down to a proper consistence

This has been incorporated in the *B P C*

MELLITE CUIVREUX (*Onguent Ægyptiac*) (*Fr*)—Copper Acetate, 1, Water, 1, Honey, 2 Boil until it assumes a red colour, and is the consistence of honey

OXYMEL DE VERDETE (*Port*)—Verdigris, 2, Vinegar, 3, Honey, 5 Boil down to a proper consistence

TOPIQUE À L'ACÉTATE DE CUIVRE (VET)—Copper Acetate, 4, Treacle, 1, Vinegar, 1, mix—*Fr*

CUPRI ACETAS—Deep green or bluish green, prismatic crystals

Solubility—1 in 15 of Water, 1 in 300 of Alcohol (90 p c), 1 in 112 of Glycerin

Medicinal Properties—Similar to the Subacetate, but more definite when required for solution in Water

Foreign Pharmacopœias—Official in *Fr*

Tests—Copper Acetate yields a bluish-green coloured solution which changes to a deep blue on the addition of Ammonia Solution in excess It should respond to the tests for Copper given under Copper Sulphate A small portion of the salt warmed with Sulphuric Acid evolves a characteristic acetous odour When warmed with Sulphuric Acid and a little Alcohol (90 p c) the odour of Ethyl Acetate is given off

The more generally occurring impurities are Arsenic, Iron, Lead and Zinc, alkalis and alkali earths Arsenic may be detected by Bettendorf's test, Lead and Zinc by boiling an aqueous solution with an excess of Sodium Hydroxide Solution, cooling, filtering, and passing Hydrogen Sulphide into the filtrate, no cloudiness or precipitate should be produced, Iron, alkalis and alkali earths by removing the Copper as Sulphide with Hydrogen Sulphide, filtering and evaporating the filtrate to dryness, no residue should remain

CUPRI SULPHAS.

COPPER SULPHATE

B P Syn—CUPRIC SULPHATE $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$, eq 247.86

FR, SULFATE DE CUIVRE, GER, KUPFERSULFAT, ITAL, SOLFATO DI RAME, SPAN, SULFATO CUPRICO

Large, translucent, blue, triclinic crystals, possessing a metallic and astringent taste. It slowly effloresces in dry air.

It may be obtained by the action of Water and Sulphuric Acid on Copper or by dissolving Copper Oxide in Sulphuric Acid.

When rendered anhydrous by heating the powder is white.

Solubility—1 in $3\frac{1}{2}$ of Water, 2 in 1 of Water (at 212°F), insoluble in Alcohol (90 p c), 1 in $2\frac{1}{2}$ of Glycerin.

Will not dissolve 1 in 275, as stated by some authorities—*P J* '02, 1 558

Medicinal Properties—Astringent, prompt emetic, eschaotie. Recommended in chronic diarrhoea, especially that of phthisis. Externally, as a styptic for bleeding surfaces and a local stimulant to ulcers, as an escharotic for warts, etc. For lotions, in proportions from 2 to 4 grains to 1 oz, also 8 grains to 1 oz for prurigo. As an astringent, **injection** to diminish excessive secretion from mucous membranes, especially in leucorrhœa and gonorrhœa. For urethral **injections**, 1 to 4 grains in an oz of Water. It is also used 1 to 2 grains to 1 oz, in granular conjunctivitis and various affections of the eyes when astringent applications are required.

An antidote in Phosphorus poisoning—3 grains every few minutes till vomiting is produced—*Mitchell Bruce*

Copper Sulphate 10 grains, Tincture of Opium 60 minims, Water 4 fl oz. This was used as a rectal injection in a bad case of dysentery—*L* '89, 11 739

Recommendation of the Departmental Committee to inquire into the use of preservatives and colouring matter in food. The use of Copper salts in the so-called greening of preserved food be prohibited—*L* '01, 11 1683, *J S C I* '01, 1228

Dose.—As $\frac{1}{4}$ to 2 grains = 0.016 to 0.13 gramme, as an emetic, 5 to 10 grains = 0.32 to 0.64 gramme

Ph Ger maximum single dose, 1.0 gramme

Prescribing Notes—*Best given in form of pill. A good pill is prepared by adding 1 part of Pulvis Tragacanthæ Compositus, and Dispensing Syrup, q s, varnish to the pill.*

Incompatibles—Alkalis and their Carbonates, Lime Water, Iodides, and most vegetable astringents.

Not Official—Guttæ Cupri Sulphatis, Cupri Oleas, Unguentum Cupri Oleatis, Lapis Divinus (Cuprum Aluminatum), Fehling's Solution, Pavy's Solution, Cupraiol, and Cupri Sulphocarbolas.

Antidotes—In case of poisoning by Copper or White of Egg is the best antidote, the stomach should be emptied, and demulcent drinks given followed by internally or Morphine hypodermically, and Linseed Meal poultices applied to the abdomen.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital (Solfato di Rame), Jap, Mex (Sulfato de Cobre), Norw, Port, Russ, Span, Swed, Swiss and U S. Ger and Swiss have also a crude sulphate.

Tests—Copper Sulphate dissolves in Water, yielding a solution which has an acid reaction towards Litmus paper, and which with Hydrogen Sulphide produces a brownish-black precipitate insoluble in Diluted Hydrochloric Acid and in Ammonium Hydrosulphide or Potassium or Sodium Hydroxide Solution, but soluble in Nitric Acid. A similar precipitate is yielded by Ammonium Hydrosulphide Solution. Ammonia Solution added drop by drop to an aqueous solution produces at first a pale blue precipitate which dissolves in an excess of the reagent yielding an intensely blue-coloured solution, Potassium or Sodium Hydroxide Solution gives a somewhat similar precipitate, which becomes brownish-black on boiling. The light blue precipitate is soluble in a very large excess of concentrated Potassium or Sodium Hydroxide Solution, but the presence of a soluble Tartrate prevents the precipitation, a deep blue liquid being produced which readily undergoes reduction to red Cuprous Oxide on boiling with Glucose and some Sugars. Potassium Hydroxide Solution produces no precipitate, but only a deep blue coloured solution, when a fixed organic acid is present. Ammonium Carbonate Solution yields a greenish-blue precipitate soluble to a deep blue solution in an excess of Ammonia Solution. Potassium Ferrocyanide Solution yields a reddish brown precipitate insoluble in dilute mineral acids, but decomposed by Potassium or Sodium Hydroxide Solution. A strip of bright metallic Iron immersed in a solution acquires a reddish coating of metallic Copper. A solution of Copper Sulphate acidified with diluted Hydrochloric Acid yields on the addition of Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid.

The more generally occurring impurities are Arsenic, Aluminium, Iron, Lead, and Zinc. Arsenic may be detected by the Bettendorf's test, Iron, Aluminium, and Zinc by removing the Copper from an acidified solution with Hydrogen Sulphide, adding Ammonia Solution to one portion and evaporating the other to dryness, in the former case no turbidity should be produced, in the latter no residue should remain. Their presence may also be ascertained by removing the Copper as oxide by boiling with Potassium or Sodium Hydroxide Solution and examining the filtrate (after acidification with Acetic Acid) by the time-limit test for heavy metals.

Heat—When heated, Copper Sulphate loses its Water of crystallisation, two molecules are lost at 30°C (86°F) with the formation of a pale blue amorphous powder, another two are lost at 100°C (212°F), and the last molecule is given up at a temperature of 200°C (392°F), leaving an anhydrous powder weighing 68.9 p.c. of its original weight. At a still higher temperature Sulphur Dioxide and Oxygen are given off, leaving a residue of Cupric Oxide, *U S P*.

Hydrogen Sulphide—If to an aqueous solution of 0.5 gramme of Copper Sulphate an excess of Hydrogen Sulphide T.S. be added and the precipitate produced filtered off, the colourless filtrate after the addition of T.S. of Ammonia should remain colourless, and after evaporation should not leave a weighable residue, *P G*.

If Hydrogen Sulphide gas be passed through 10 c.c. of an aqueous solution (1-20) of the salt, to which 1 c.c. of diluted Hydrochloric Acid has been previously added, until precipitation of Copper Sulphide is complete, one half of the colourless filtrate should not be coloured or rendered turbid on the addition of T.S. of Ammonia, nor should the other half yield a weighable residue when evaporated and ignited, *U S P*.

Time-limit Test—An aqueous solution of the salt (1-20) when boiled with excess of TS of Sodium Hydroxide until the Copper Oxide is completely dissolved and then filtered, a colourless filtrate should be obtained which, Acetic Acid, should not respond to the time-limit test for heavy metals, *U S P*

Not Official

GUTTÆ CUPRI SULPHATIS—Copper Sulphate, 2 grains, Water, 1 fl oz —*London Ophthalmic* The strength in use at the principal hospitals

CUPRI OLEAS—Green, oleaginous solid, insoluble in Water, soluble in Ether An excellent antiseptic and antiparasitic agent When diluted it is especially useful in ringworm

UNGUENTUM CUPRI OLEATIS—Copper Oleate, 1, Lard, 4, melt together, and stir till cold Useful in ringworm, hard and horny warts, corns and bunions —*B M J '84*, 11 752

1 to 7 of Soft Paraffin (*London*), 1 to 7 of Lard (*University*), 1 to 9 of Lard (*B P C*)

LAPIS DIVINUS CUPRUM ALUMINATUM—Copper Sulphate, Potassium Nitrate, and Alum, of each equal parts, in powder, fused in a glazed earthen crucible, powdered Camphor, to the extent of $\frac{1}{4}$ th part of the whole, being added near the end of the process When cold, break in pieces and keep in a close stoppered bottle An eye-wash may be made by dissolving 2 grains in 1 oz of distilled Water

Foreign Pharmacopœias—Official in Fr (*Pierre Divine*), Ger, Hung, Jap, Russ and Swiss Not in the others

FEHLING'S SOLUTION—*See Appendix*

PAVY'S SOLUTION—Crystallised Copper Sulphate, 34.65 grammes, Rochelle Salt, 170 grammes, Potassium Hydroxide, 170 grammes, Water, to 1000 cc

When 120 cc of this Solution are mixed with 400 cc of Ammonia (sp gr 0.880) and diluted to 1000 cc, then 10 cc may be taken as equivalent to 0.005 gramme of Glucose

The method is well adapted for the examination of Diabetic Urine and Milk, also mixtures of Milk and Cane Sugars, and certainly has the advantage over the ordinary Fehling method by its definite end reaction

CUPRARGOL (Copper and Silver Albuminate)—A greyish-white powder, soluble in Water Has been used in 1 to 5 p c solution in conjunctivitis

CUPRI SULPHOCARBOLAS (Copper Aseptol)—Green rhombic prisms or light green needle-shaped crystals Soluble in Water and in Alcohol (90 p c) Hemostatic As an antiseptic, $\frac{1}{2}$ to 1 p c solution As an injection in gonorrhœa, $\frac{1}{2}$ to 1 $\frac{1}{2}$ p c solution

Not Official

CURARA—WOORARA.

A powerful poison, stated to be obtained from various species of *Strychnos* and other plants, and used by the Indians in the northern part of South America for arming the points of their arrows A brownish black, shining, brittle, resinous mass, almost wholly soluble in Water, sparingly soluble in Absolute Alcohol Different samples may vary very much in strength, and no doubt also in general composition so that the dose of every parcel has to be arrived at by experiment It is only used hypodermically, and the solution has generally been of the strength of 1 grain in 100 minims

An alkaloid Curarine has been obtained from Curara, and although commercial, is somewhat difficult to obtain

Arrow Poisons Their history, sources, and constituents —(*Stockman*) *P J '98*, 11 548, 585

Medicinal Properties—It has been used in the treatment of Strychnine poisoning, hydrophobia, chorea, and tetanus

In the convulsions of chorea, and to prevent painful spasms in moving wounded persons (*B M J* '04, 11 1642) Has an influence in diminishing the severity of the tonic and clonic tetanic spasms (*L* '05, 1 991), and should be injected hypodermically morning and evening, commencing with $\frac{1}{2}$ grain and gradually increasing the dose according to the severity of the spasms to 1 grain also 20 to 30 c c of antitetanic serum injected under the skin of the abdomen and repeated daily for a week or ten days

Dose— $\frac{1}{12}$ to $\frac{1}{2}$ grain = 0.005 to 0.032 gramme, but should be used with great care

Foreign Pharmacopœias—Official in Mex (Curaro) Not in the others

Descriptive Notes—Curare is usually imported from Venezuela in the form of a blackish extract contained in small gourds about $2\frac{1}{2}$ inches in diameter A fragment of the extract placed in a drop of Alcohol on a microscopic slide shows a brownish fluid copiously studded with quadrilateral prisms (supposed to be Curarine, which forms four sided prisms) and an abundance of minute particles of a yellowish tint which consist of Calcium Oxalate The principal ingredient in Curare is the bark of *Strychnos toxifera*, but that of other species is also used in different districts, thus in British Guiana the bark of *S. toxifera*, Schomb, *S. Schomburgkii*, Kl, and *S. cogens*, Benth, are used in the Curare of the Macusi, Oreocua and Wapishana tribes, that of *S. Gubleri*, Planch, by the Moquitari and Puaroa Indians, between the Orinoco and Rio Negro, and that of *S. Castelnociana*, Wedd, by the Ticuna, Pebia Yagua and Oregona Indians in the districts of the Upper Amazon, that of *S. Orevarum*, Planch, by the Tio and Roucouyenne Indians of French Guiana It is obvious, therefore, that Curare is an extract of uncertain composition as regards the species of *Strychnos* employed, and the more so that different ingredients are added to the extract by different tribes A preparation made in this country from the bark of *S. toxifera* imported from British Guiana would, therefore, be far more reliable Curare is said to have been used successfully in hydrophobia

INJECTIO CURARÆ HYPODERMICA.—Curare, 1^g, make it into a paste with Distilled Water recently boiled and cooled, transfer to a funnel plugged with absorbent Wool, and gradually pour upon it Distilled Water until 10 is obtained —*B P C*

B P C Formulary 1894 gave the strength as 1 grain in 12 minims

Dose—1 to 6 minims = 0.06 to 0.36 c c

CUSPARIÆ CORTEX.

CUSPARIA BARK

FR, ANGUSTURE VRAIE, GER, ANGOSTURARINDE, ITAL, CORTECCIA DE ANGUSTURA, SPAN, CORTEZA DE ANGOSTURA

The dried Bark of *Cusparia febrifuga*, DC

The alkaloids, **Cusparine**, **Cusparadine**, **Galipeine**, and **Galipidine** have been extracted from Cusparia Bark The bitter principle **Angosturin**, to which the bitterness of the bark has been assumed to be due has also been isolated, and a small quantity of a volatile Oil **Cusparine Sulphate** and **Hydrochloride** are slightly soluble in Water, the Acetate and Tartrate much more so —*P J* (3) xiv 423

Contains about 1.5 p c of ethereal oil —*J C S Abs* '98, 1 87

Medicinal Properties—An aromatic bitter tonic In South America it is given as an antiperiodic for malarial fever

Prescribing Notes—Given in the form of the Infusion or the Concentrated Infusion, generally combined with Aromatics to prevent nausea

Official Preparations—Infusum Cuspariæ and Liquor Cuspariæ Concentratus

Foreign Pharmacopœias—Official in Mex (Angostura Verdadera) and Port Not in the others

Descriptive Notes.—The bark of *Cusparia febrifuga* varies somewhat in appearance and size, the outer surface being in some pieces smooth and hard, and in others soft and spongy. The pieces are usually curved, 2 to 4 or more inches long (4 to 5, *BP*), about $1\frac{1}{2}$ inch in thickness and 1 inch or more in diameter. Its distinctive characters are the short, resinous, brown fracture, laminated inner surface, characteristic odour and flavour, in addition to its bitterness.

The character mentioned in the *BP* of numerous white points on the transverse fracture is also found in *Nux Vomica* bark, which was at one time substituted for it in commerce. The latter, however, has a paler fracture with a definite paler line. The corky portion, is not laminated, and is purely bitter without any special flavour. In powder *Cusparia* Bark may be recognised by the presence of numerous oil cells, by the acicular as well as single, oblique prismatic crystals of Calcium Oxalate, and by having very thick-walled bast fibres, which are coloured canary-yellow by Caustic Potash.

Tests.—*Cusparia* Bark yields about 8 p.c. of ash, and 10 p.c. is rarely exceeded.

Preparations.

INFUSUM CUSPARIÆ INFUSION OF CUSPARIA.

Cusparia Bark, in No. 20 powder, 1, Distilled Water, boiling, 20; infuse for fifteen minutes, strain (1 in 20)

Dose.—1 to 2 fl oz = 28.4 to 56.8 c.c.

Incompatibles—Mineral Acids, Ferric Chloride, and other metallic salts.

LIQUOR CUSPARIÆ CONCENTRATUS. CONCENTRATED SOLUTION OF CUSPARIA

10 of *Cusparia* Bark, in No. 40 powder, percolated with Alcohol (20 p.c.), to yield 20 (1 in 2)

Dose.— $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c.c.

Tests—Liquor *Cuspariæ* Concentratus has a specific gravity of 1.010 to 1.020, contains about 10.0 p.c. w/v of total solids and about 20.0 p.c. w/v of Absolute Alcohol.

Infusum Cuspariæ Concentratum—*Cusparia* Bark, in No. 40 powder 40, Alcohol (90 p.c.), 25, Distilled Water (1 in 1000), q.s. to make 100. Prepare by maceration. —*Farr and Wright, P J '06, 1 165 and '07, 1 621, C D '06, 1 252, Y B I', '07 219*

This appears in the *BP C*

CUSSO.

KOUSSO

FR, COUSSO, GER, KOSOBLOUTHEN, ITAL, KOUSSO, SPAN, FLOR DE COUSSO

The dried panicles of pistillate Flowers of *Brayera anthelmintica*, Kunth (of *Hagenia abyssinica* (Bruce) Gmelin, *USP*)

Obtained from Abyssinia

Medicinal Properties—Anthelmintic Especially useful for the different kinds of tapeworm Should be followed by a purgative to expel the dead worm

Dose— $\frac{1}{4}$ to $\frac{1}{2}$ oz = 7 1 to 14 2 grammes

Prescribing Notes—*The Flowers, in coarse powder, are mixed with half a pint of warm Water, allowed to stand for fifteen minutes, stirred up (not strained), and taken in 2 or 3 draughts at short intervals It should be taken in the morning on an empty stomach, the bowels having previously acted After three or four hours a brisk purgative should be administered On account of its liability to produce nausea a little Lemonade may be taken afterwards*

Foreign Pharmacopœias—Official in Austr (Koso), Belg, Fr and Port (Coussou), Ital (Koussou), Ger, Jap and Swiss (Flores Koso), Hung (Kusso), Mex (Cuso), Norw and Swed, Russ (Flores Kusso), Span (Couso), US (Cusso)

The Infusion is official in Belg 2 in 15, Fr (Apozeme de Coussou) about 1 in 8, Span (Inf de Couso), 1 in about 16 $\frac{1}{2}$, *BP C* 6 in 100

Descriptive Notes—Cusso is generally imported in rolls about 1 $\frac{1}{2}$ to 2 feet long (3 to 6 dm) and 2 $\frac{1}{4}$ to 3 inches in diameter, each consisting of the large paniced inflorescence of female flowers The flowers are shortly stalked, pubescent, and consist of the two rounded bracts, a calyx of 5 outer, rigid, purple-veined sepals, and 5 inner, smaller, incurved, shrivelled segments The 5 caducous white petals as well as the abortive stamens are usually absent in the drug, there are two carpels, and the style with two broad hairy stigmas is exerted The large stems should be rejected (*USP*), and also samples which have lost their reddish tint and become brownish, indicating age and deterioration The male flowers, which are excluded by the *BP*, *USP* and *PG*, have a greenish tint, and the outer sepals are not enlarged The powder is characterised by the thick-walled unicellular hairs, the numerous rosette crystals, as well as simple and segmented capitulate short stalked glands (*Koch*) It should not contain pollen grains, nor fragments of vessels more than 0.002 mm in diameter (*PG*), but *Koch* states that pollen grains do not occur in it

Not Official

~~CYDONIUM~~

~~QUINCE SEED~~

~~The Seeds of *Pyrus Cydonia*~~

~~Their coriaceous envelope abounds in Mucilage~~

~~**Medicinal Properties**—Demulcent The decoction is used externally for cracks in the skin A nice adjunct to eye lotions in cases of irritation and inflammation~~

Foreign Pharmacopœias—Official in Belg, Port (Maimelo), Mex (Membrillo), and Swiss Not in the others The fruit is official in Fr (Coing)

DECOCTUM CYDONII—Quince Seed, 1, Distilled Water, 80 Boil for ten minutes, and strain

This has been incorporated in the *BPC*

MUCILAGO CYDONII—1 of Quince Seed and 25 of Water, by cold maceration Was official in Austrian Ph (1889), and is the strength now incorporated in the *BPC*

Fr, 1 in 10, Port, 1 in 100

Not Official.

CYLLIN.

Jeyes' Cyllin (medical) is a non-toxic antiseptic, it contains 50 p.c. of a new series of oxidised Hydriocarbons, free from Phenol and its homologues, emulsified with neutral Tar Oil Its Carboic Acid coefficient for *Bacillus Typhosus* is 20 —*L* '07, 1 33

Preparations for internal administration are Cyllin Capsules, Cyllin Pastilles and Cyllin Syrup

The dressings, Cyllin Lint, Gauze and Wool, each contain 10 p.c. of Cyllin

Two Gelatin plates were inoculated with *Staphylococcus pyogenes aureus*, one was medicated with Cyllin inhalant, and the other was unmedicated The former showed no sign of growth, the latter a copious and normal growth —*L* '05, 1 988 A new use of the inhalant is thus indicated in combating the symbiotic action of the pus microbe in pulmonary tuberculosis

3-minim doses given (*L* '05, 1 1148), in the form of a keratin coated capsule, every second hour if necessary in the treatment of sprue, but rarely more than 8 a day are required Best time to administer it is after food

Attention has been called (*B M J* '04, 1 1119) to its value as a tubercle destroyer it being claimed to be non-toxic and about twenty times as powerful as Carbolic Acid, and hence its trial in cases of tuberculosis —*L* '05, 1 377

Not Official

CYNOGLOSSUM

The Root of *Cynoglossum officinale*, *L* It contains an amorphous alkaloid **Cynoglossine**

Medicinal Properties—Has been used as a demulcent and sedative

Foreign Pharmacopœias—Official in Dan, Fr, Mex, Norw, Port and Span Not in

Pulvis Cynoglossi Compositus is official in Dan, Fr and Norw, **Pilulas Cynoglossi** in Dan, **Pildoras de Cynoglossa** in Mex, and Span

Fr, **Pilules de Cynoglosse Opiacees**, each pill contains 0.02 gramme ($\frac{1}{50}$ grain) of Extract of Opium, and 0.02 gramme of Powdered Henbane Seeds

Not Official

CYPRIPEDIUM

The Rhizome and Roots of *Cypripedium florum* Mill, and of *C. pubescens* Salisb, are official in U.S. Pharmacopœia known as Lady's Slipper

Cypripedin is the dried extract of the above, and has been recommended in hypochondriasis, chorea and epilepsy

The eclectic remedy 'Cypripedin' which is stated to be prepared by precipitating a concentrated tincture of the root with Water, is complex in composition and stated to have no claim to the name given it

It may be used in doses of 1 to 5 grains = 0.065 to 0.32 gramme

A 1 in 1 Fluid Extract (Alcohol 48.9 p.c.) is official in the U.S., average dose 15 minims = 0.9 cc

Not Official

DAMIANA

The Leaves of one or more species of *Turnera*, from Mexico and California. Contains a bitter substance, resins, and a volatile oil.

Medicinal Properties—Tonic, diuretic, and aphrodisiac.

Prescribing Notes—Frequently given in the form of pill, the Hard Extract makes a good pill with a small quantity of Alcohol (90 p.c.), the Soft Extract is best hardened with the powdered Leaves. The Liquid Extract is given in capsules.

Foreign Pharmacopœias—Official in Mex.

Descriptive Notes—Damiana occurs in commerce in two or three different varieties. The kind which is considered the best is known as *Helmichis Damiana*, and is derived from *Turnera aphrodisiaca*, L. Ward (nat. ord. *Turneraceæ*) and is a native of California and Mexico. The leaves, of a light green colour, are wedge shaped, usually less than an inch in length (10 to 25 mm) and about $\frac{1}{2}$ inch (5 to 10 mm) in diameter in the broadest part, with about 3 to 6 coarsely crenate teeth on either side besides the terminal one. The taste recalls that of figs, but is aromatic and slightly bitter. The plant is considered by some botanists to be a variety of *T. diffusa*, Willd. It has reddish stems and thinner, smoother and less hairy, greener leaves, not greyish green as in the type. The leaves of another species, possibly *T. microphylla*, Desv., are sometimes substituted for it. They are smaller, more hairy, with hairy stems which are not reddish. The leaves of *Aplopappus discordeus*, DC., nat. ord. *Compositæ*, are occasionally offered as Damiana. They have fewer, more distant, serrate teeth, usually three on either side, and composite flowers with hairy pappus, usually mixed with the leaves.

EXTRACTUM DAMIANÆ LIQUIDUM—Damiana leaves exhausted with Alcohol (60 p.c.), 1 of fluid represents 1 of the drug.

This has been incorporated in the B.P.C.

Dose—30 to 60 minims = 1.8 to 3.6 cc.

EXTRACTUM DAMIANÆ—The above evaporated to a soft extract.

Dose—5 to 10 grains = 0.32 to 0.65 gramme.

This has been incorporated in the B.P.C.

MISTURA DAMIANÆ COMPOSITA—Sodium Hypophosphite, 5 grains, Calcium Hypophosphite, 5 grains, Liquid Extract of Damiana, $\frac{1}{2}$ fl. drm., Liquid Extract of Nux Vomica, 2 minims, Chloroform Water, to 2 drms.—*Martindale*.

This has been incorporated in the B.P.C.

PILULA DAMIANÆ COMPOSITA—Extract of Damiana, 2 grains, Phosphorus, $\frac{1}{10}$ grain, Extract of Nux Vomica, $\frac{1}{2}$ grain.—*Martindale*.

Extract of Damiana, 2 grains, Extract of Nux Vomica, $\frac{1}{10}$ grain, Phosphorated Suet (10 p.c.), $\frac{1}{10}$ grain, mix quickly these three with about $\frac{1}{2}$ minim of Chloroform and add $\frac{1}{10}$ grain of Compound Tragacanth Powder and Mucilage of Acacia q.s.—B.P.C.

DIGITALIS FOLIA.

DIGITALIS LEAVES

FR., DIGITALE, GER., FINGERHUTBLÄTTER, ITAL., DIGITALE, SPAN., HOJA DE DIGITAL.

The dried leaves of *Digitalis purpurea*, L. Collected from plants commencing to flower. The U.S.P. specifies that the leaves should be collected from plants of the second year's growth.

Medicinal Properties—Cardiac and circulatory stimulant and tonic, increases the strength and efficiency of the cardiac contractions, and reduces the pulse rate without diminishing tension. Specially useful in mitral and tricuspid lesions with loss of compensation, in cardiac insufficiency from whatever cause, with irregular and rapid action and low arterial tension, not indicated in purely aortic cases. Of great value as a cardiac stimulant in acute pneumonia, useful in pulmonary hæmorrhage due to mitral disease. Diuretic, useful in cardiac dropsy, also in renal dropsy when acute or when due to failure of a hypertrophied heart.

It is cumulative in action, and requires watchfulness. Its continued use deranges the alimentary system, therefore, after it has been taken for eight or ten days it should be left off for three or four days and then recommenced. According to Lauder Brunton, Digitalis is distinctly dangerous in advanced fatty degeneration of the heart, he also thinks it harmful in advanced Bright's disease. For a comparison with *Strophanthus* see under *Strophantha Semina*.

According to Kihani, the seeds of *Digitalis purpurea* contain Digitalinum verum, and Digitonin, the leaves contain Digitoxin, but neither of the other two. Preparation of Digitalin also described—*JCS Abs* '96, 1 58, 59, 180, '97, 1 95, *PJ* '95, 11 29, 120, '96, 11 289.

Treatment of pneumonia by Digitalis—*B M J E* '95, 11 82, '96, 11 76, '97, 1 15.

Digitoxin in doses of $\frac{1}{2}$ milligramme = $\frac{1}{288}$ grain—*B M J E* '97, 1 81.

Best administered as tincture or dried leaf in pill form, one advantage being combination of active principles whereby overaction is obviated—*L* '02, 1 673.

Untoward effects of 20 to 24 minims of the tincture taken daily for five days in a weakly overgrown boy of 10 years—*B M J E* '02, 1 1068.

The administration of Digitalis particularly deprecated in unduly high blood pressure—*B M J* '99, 1 85.

In the treatment of the rapid heart of influenza, where there is cardiac dilatation that lasts some weeks. Intermittent administration best. 10 to 20 minims of Tincture, $\frac{1}{2}$ drm of the Infusion or 1 grain Powdered Leaves, three daily for three days with intervals of three or more days during which the drug is withheld, or 2 to 4 granules of Nativelle's Digitalin containing $\frac{1}{16}$ milligramme once in twenty-four hours for two days with intervals of at least three days—*L* '99, 11 1079.

Some remarks upon Digitalis treatment in chronic disorders of the circulation, and especially upon the continuous use of Digitalis—*P* lxiv 385.

An experimental investigation into the treatment of Digitalis poisoning. Nitroglycerin, besides possessing a greater antagonistic action to Digitalis than any other known drug, is also relatively non-toxic, and for the reduction of internal tension Nitroglycerin or an ally is the best remedy, but with a low blood pressure these substances are useless—*B M J* '99, 11 1265.

Influence on the heart muscle when administered for a long time—*T G* '97, 800.

Although this is the most powerful remedy in mitral disease which we possess, in cases of pure aortic regurgitation (*B M J* '06, 1 1462), it is not only a dangerous drug but is actually a poison. It acts as a diuretic in cardiac dropsy (*B M J* '06, 1 10). The diuresis produced by the ordinary doses of the pharmacopœial preparations of the drug does not make its appearance before the third or fourth day of administration. To be useful in cardiac dropsy it must be given in full doses.

The pharmacological action of Digitalis, *Strophanthus*, and Squill is fully considered—*B M J Supplement*, '06, 11 13, *PJ* '06, 11 28.

Dose—In powder, $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme.

Ph Ger maximum single dose, 0.2 gramme, maximum daily dose, 1.0 gramme

Prescribing Notes—*The fresh Infusion is preferred by some to the Tincture. The powdered leaf is ordered in Pills with other ingredients. Ferrous Sulphate is not uncommonly prescribed with the fluid preparation of Digitalis, with a resulting blackening from the tannin of Digitalis where this is an objection it can be prevented by the addition of Citric Acid. 6 grains of Citric Acid are sufficient for 12 grains of Ferrous Sulphate, or the powdered drugs can be given in pills.*

Incompatibles—Ferrous Sulphate, Tincture of Ferric Chloride, preparations of Cinchona, and Lead Acetate

Official Preparations—Infusum Digitalis and Tinctura Digitalis

Not Official—Fluidextractum Digitalis, Infusum Digitalis Concentratum, Pilula Digitalis Composita, Pilula Digitalis et Hydrargyri Composita, Pilula Digitalis et Opi Composita, Succus Digitalis, Syrupus Digitalis, Vinum Digitalis Compositum, and Digitalin (various)

Antidotes—In case of an overdose, a recumbent posture is of paramount importance, and after the stomach has been emptied, 20 grains of Tannic Acid in hot Water given frequently, or hot strong Tea or Coffee, stimulants and warmth should be employed

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port (Dedaloira), Russ, Span (Digital), Swed, Swiss and U.S.

The Brussels Conference agreed to use the leaf of the second year, and the powdered drug to be used entire

Tests—The ash of Digitalis varies from 7 to 10 p.c., and should not exceed the latter figure. In the present state of our knowledge of the subject a chemical method of determination is of doubtful utility, and it is generally conceded that a physiological method of standardisation leaves much to be desired.

The more or less definite principles contained in Digitalis may be arranged as follows under the names applied to them by Schmiedeberg:

- (a) **Digitonin**.—A crystallisable body resembling Saponin, constituting the larger part of the glucosidal constituents. It softens at 225° C (437° F), and melts completely at 235° C (455° F). The aqueous solution is levorotatory, and is precipitated by Tannin, Ammoniacal Lead Acetate, and Barium Hydroxide Solution. Soluble in Water, insoluble in cold Alcohol, Ether, Benzol, or Chloroform. It has none of the physiological action peculiar to Digitalis, and in other respects is directly injurious.
- (b) **Digitalein**.—An amorphous glucoside (possibly a mixture). Soluble in Water and in Alcohol, insoluble in Ether or Chloroform. Its action on the heart is non-cumulative, and it causes no irritation when subcutaneously injected.
- (c) **Digitalin**.—A white amorphous powder or soft white grains which remains unchanged when heated to 200° C (392° F), begins to aggregate at 210° C (410° F), and melts about 217° C (422° F). It is soluble in Alcohol, almost insoluble in Water, sparingly soluble in Ether or Chloroform. It dissolves in concentrated Hydrochloric or Sulphuric Acid with golden yellow colour, the colour in the latter instance changing rapidly to a blood red. On adding to the solution whilst still yellow a drop of either Nitric Acid, Ferric Chloride, or Bromine Water, a brilliant purple coloration is produced.

Possesses in a high degree the medicinal action of Digitalis

- (d) **Digitoxin**.—Pearly plates or needles melting at 240° C (464° F). *Fr Codex* (1908) gives 243° C (469° F). Easily soluble in Alcohol, slowly in Chloroform, very sparingly in Ether, quite insoluble in Water. It

does not give the colour reaction of Digitalin with strong Sulphuric Acid. It yields a yellow or greenish coloration when warmed with strong Hydrochloric Acid. The *French Codex* gives the following test. If weighed quantities of about 0.005 gramme of Ferric Sulphate be dissolved separately in 2 c.c. of Glacial Acetic Acid and in 2 c.c. of concentrated Sulphuric Acid, then if the Acetic Acid Solution of the Iron salt, previously mixed with a trace of Digitoxin, be poured gently on to the surface of the Sulphuric Acid containing the Iron salt in such a manner that the two liquids do not mix, at the point of contact of the two liquids there is developed a brown coloured zone, the colour changing to green and then to indigo-blue and after half an hour the Acetic Acid is coloured entirely blue. This characteristic reaction for Digitoxin was given in the 17th edition of the *Companion* as follows. When dissolved in Acetic Acid to which has been added 1 p.c. of a solution containing 5 p.c. of Ferric Sulphate, and Sulphuric Acid containing the same quantity of Ferric Sulphate is poured into the tube so as to form a layer beneath it, a blue colour is gradually developed in the Acetic Acid, whilst the Sulphuric Acid remains colourless, this coloration in the Acetic Acid is not produced by any other of these compounds.

The most toxic of all the constituents, but uncertain, cumulative, and dangerous in its action.

- (e) **Digitalin**—A crystalline body, physiologically inert, difficultly soluble in Water, more readily in Alcohol, insoluble in Ether or Chloroform.

It dissolves in concentrated Sulphuric Acid with brownish-yellow colour, which becomes purple-red on exposure to air, the addition of Water turning the colour to green. It is insoluble in Hydrochloric Acid, but dissolves in Nitric Acid without coloration. It does not reduce Fehling's Solution until after boiling with dilute acid.

Descriptive Notes—The Digitalis leaves of commerce are probably collected, not only from plants in flower, but from plants of the first year's growth and from shoots formed laterally from the second year's plant after flowering. The leaves accordingly differ in size and shape. The root leaves are broadly ovate, lanceolate, and crenate-serrate, and much larger and wider at the base than the stem leaves, which have shorter stalks and are gradually less ovate and more lanceolate and smaller from the base to the top of the leafy stem. The dimensions given in the *B.P.*, viz., 10 to 30 cm (4 to 12 inches) long, and 12½ to 15 cm (5 or 6 inches) broad, cover all the above forms of leaf. The leaves have a bitter taste, but no distinct odour. The features that distinguish Foxglove leaves from other similar leaves are the lower veins, which are placed at a very acute angle to the base of the mid-rib and are decurrent into the petiole, the faintly areolate ~~the~~ *B.P.* upper surface, and the paler under surface, densely pubescent with short hairs, and reticulated with prominent small veins. Other leaves occasionally found mixed with those of Foxglove differ in the venation and in the character of the hairs, which in Digitalis are of two kinds, viz., 1 to 5 celled simple hairs (usually 3 celled, *B.P.*), with occasionally a glandular head, and short 1 to 2 celled hairs, with a single or a twin gland at the apex.

The powdered leaves are distinguished by the absence of raphides and stone cells, by sinuous epidermal cells with small stomata, the 1-5-celled hairs with a more or less warty surface, and short glandular hairs.

Digitalis leaves soon lose their activity unless kept quite dry, they can be obtained in commerce with the midrib removed and packed dry in hermetically sealed bottles. The tincture would probably be more effective if prepared from the fresh leaves with rectified spirit and then heated to boiling point to prevent changes arising from the action of natural ferments in the leaves, since this takes place in the leaves in the presence of the 12 p c of moisture which they absorb after drying, if not enclosed in airtight vessels as soon as dried. The *P G* points out the triangular character of the petiole, and that the upper stem leaves may be sessile. The leaves should be obtained from wild plants. The *USP* directs leaves collected from the second year growth at the commencement of flowering, and that stone cells, star shaped hairs, and Calcium Oxalate crystals should not be present. Continental Digitalis leaves are liable to be adulterated with many other leaves—see *Apotheker Zeitung*, xxi pp 242, 252, 267, 276, where the distinctive features of the probable adulterants are given.

Preparations

INFUSUM DIGITALIS INFUSION OF DIGITALIS

Digitalis Leaves, in No 20 powder, 60 grains, boiling Distilled Water, 20 fl oz. Infuse fifteen minutes, strain.

Dose—2 to 4 fl drm = 7.1 to 14.2 c c

1 fl oz represents 3 grains of Leaves

Foreign Pharmacopœias—Official in Mex, Port and Spain, 1 in 200, Ital and Swed, 1 in 100, U S, with Cinnamon, 3 in 200. Not in the others.

Infusum Digitalis Concentratum—Digitalis Leaves, in No 20 powder, 55, Alcohol (90 p c), 20, Dilute Chloroform Water (1 in 1000), q s to make 100. Prepare by macero expression. Dose, 15 to 30 minims—*Farr and Wright, P J* '06, 1 165, and '07, 1 622, *C D* '06, 1 252, *Y B P*, 1907, 249.

This appears in the *B P C*

TINCTURA DIGITALIS TINCTURE OF DIGITALIS

2½ of Digitalis Leaves, percolated with Alcohol (60 p c) to yield 20 (1 in 8)

Dose—5 to 15 minims = 0.3 to 0.9 c c

Ph Ger maximum single dose, 1.5 grammes, maximum daily dose, 5.0 grammes, of the 1 and 10 Tincture.

Larger doses are occasionally given, but, according to some observers, the results with small doses are equally good and not nearly so dangerous.

In cases of delirium tremens, 1 fl drm every three hours. Two or even three fl drm in cases carefully watched—*P J* xxvii 373.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Russ, Swiss and U S, 1 in 10. Ger, Ital, Jap, Norw, Spain and Swed, 1 and 10, Hung and Port, 1 in 5. Also Port and Spain, *Fresh Leaves*, 1 Spirit, Spain, with Ether, 1 dried Leaves in 5, Dan and Port, with Spirit of Ether, 1 dried Leaves in 10, Mex, Seeds 1 in 5, also *Infusum* Tincture 1 and 5. All by weight except U S.

The *Brussels Conference* agreed to a strength of 10 p c prepared by percolation with Alcohol (70 p c). Belg, Dan, Fr, Ital, Jap, Russ and Swiss adopt this.

Tests—Tincture of Digitalis possesses a specific gravity of 0.930 to 0.935, contains from 3 to 4 p c w/v of total solids and about 55.0 p c w/v of Absolute Alcohol.

A suggestion for the preparation of a fat-free Tincture (*A J P* '99, 332), by exhausting the leaves with purified Petroleum Benzin previous to the preparation of a Tincture by the official process. The resulting tincture is claimed to be less nauseating than the ordinary tincture, and to be more rapidly absorbed. Abscesses never followed its hypodermic use, whilst the official Tincture almost invariably causes pain, swelling, and abscess formation.

A fermentation test has been suggested (*C D* '02, 1 456) to prove the activity of *Digitalis* leaves. A weighed quantity of 20 grains of Amygdalin is dissolved in 1 fl oz of Water at 30° C (98° F) in a wide-mouthed bottle and set aside as a control specimen. A similar quantity of Amygdalin, together with 60 grains of powdered *Digitalis* leaves, is mixed in another bottle. At the end of eight hours the plain Amygdalin solution should show no change, but the specimen to which the *Digitalis* has been added should have developed a bitter Almond odour, and should yield a reaction for Hydrocyanic Acid when a rod moistened with Silver Nitrate solution is laid over the mouth of the bottle. The test is considered (*C D* '02, 1 509) ingenious, but inconclusive, as it does not prove the presence or absence of Digitoxin.

Not Official.

FLUIDEXTRACTUM DIGITALIS (US)—A 1 in 1 fluid extract prepared by exhausting the Leaves with Alcohol (49 p c)

Dose—1 to 2 minims = 0.06 to 0.12 c c

Foreign Pharmacopœias—Official in Dan, Mex and US. Not in the others.

Extractum Digitalis (US) is the Fluid Extract evaporated (not exceeding 50° C) to a pill consistence.

PILULA DIGITALIS COMPOSITA—*Digitalis* Leaves, in powder, 1 grain, Squill, in powder, 1 grain, Mercury Pill, 1 grain—*St Thomas's*.

This has been incorporated in the *B P C*, with the *syn* Guy's Pills, and a note that Baillie's or Gilmour's Pills contain twice as much Squill.

PILULA DIGITALIS COMPOSITA (Baillie's Pill) *Digitalis* Leaves, in powder, ½ grain, Squill, 1 grain, Mercury Pill, 2 grains, in one pill—*St George's*.

PILULA DIGITALIS ET HYDRARGYRI COMPOSITA—Mercurial Pill, 1 grain, Powdered *Digitalis*, 1 grain, Powdered Squill, 1 grain, Extract of Henbane, 2 grains—*St Bartholomew's*.

PILULA DIGITALIS ET OPII COMPOSITA (Heim's Pill)—Quinine Sulphate, 1 grain, *Digitalis*, in powder, ½ grain, Opium, in powder, ½ grain, Ipecacuanha, in powder, ½ grain, Glycerin of Tragacanth, q s—*St George's*.
Digitalis, in powder, ½ grain, Opium, in powder, ½ grain, Ipecacuanha Root, in powder, ½ grain, Quinine Sulphate, 1 grain, Syrup of Glucose, q s—*B P C*.

SUCCUS DIGITALIS—The Expressed Juice, 3, Alcohol (90 p c), 1.

This preparation may be given for a longer period than the Tincture without causing nausea.

Dose—5 to 10 minims = 0.3 to 0.6 c c

This has been incorporated in the *B P C*.

SYRUPUS DIGITALIS—Tincture of *Digitalis*, 1, Simple Syrup, 19—*Belg* and *Fr*.

VINUM DIGITALIS COMPOSITUM—Digitalis Leaves, in No 40, powder, 5, Squill, 7 5, Juniper Fruit, 75, Alcohol, 100, White Wine, 900, Potassium Acetate, 50—*Belq* and *Fr*

Alcoholic Extract 2, Water 360, Sugar 640—*Span*

DIGITALIN—Under this name four distinct varieties occur in commerce, which differ so considerably in their medicinal properties that prescribers should be careful to distinguish and specify the kind intended. All four of them are soluble in Alcohol

- 1 **Digitalin Amorphous (Homolle)**—A white or yellowish white amorphous bitter powder. Soluble in Chloroform, slightly soluble in Water. Stated to consist mainly of Digitalin with some Digitoxin. Now omitted from *Fr Codex*

Foreign Pharmacopœias—Official in Port and Span, formerly in Brit

- 2 **Digitalin Crystallised (Nativelle)**—Fine white needles, insoluble in Water, Ether or Benzene, soluble in Alcohol (90 p c) and in Chloroform. It is stated to consist almost entirely of Digitoxin, and is cumulative in its action. See Digitoxin

Foreign Pharmacopœias—Official in Fr, Mex and Span

Granules de Digitaline Cristallisée (*Fr Codex*) contains $\frac{1}{10}$ milligramme in each granule. Soluté de Digitaline Cristallisée au Millième contains one milligramme to each gramme

Digitoxin, official in *Swiss*, is a white, crystalline, odourless powder with a bitter taste, insoluble in Water, soluble in Alcohol (90 p c), and in Chloroform

- 3 **Digitalin German**—Amorphous, consists principally of Digitalin with some Digitonin and Digitalin. Readily soluble in Water, almost insoluble in Chloroform

- 4 **Digitalin Verum**—Kühn (P J (3) xxii 1061) states, with some show of reason, that the Digitalin of Schmiedeberg is the best form in which to prescribe Digitalis, and to distinguish it he applies the name Digitalin Verum. Its composition is definite, it is obtainable commercially in a sufficiently pure condition, it possesses all the medicinal activity in regard to the action of Digitalis upon the heart, it is non-cumulative in its action, the dose is $\frac{1}{4}$ milligramme ($\frac{1}{100}$ grain) every 2 or 3 hours, it is soluble about 1 in 1000 of Water, about 1 in 100 of Alcohol (50 p c). The aqueous solution froths upon being shaken, and is remarkably prone to become mouldy

Not Official

DUBOISIA MYOPOROIDES

A plant indigenous to N S Wales and Queensland, it has been classed in the order *Solanaceæ*

Ringer's experiments show that the physiological action of the extract is apparently identical with that of Atropine. Tweedy has used it as an application to the eye in all cases in which Atropine is indicated

Foreign Pharmacopœias—Official in Span

The name Duboisine represents a variable product obtained from this plant. Pseudo hyoscyamine from *Duboisia myoporoides*, R Br, isomeric with Atropine and Hyoscyamine, has been described by Merck

Foreign Pharmacopœias—Official in Mex

Duboisine Sulphate is an amorphous hygroscopic powder, soluble in Water, consisting of an indefinite mixture of Hyoscyamine and Hyoscyne Sulphates, and the Sulphates of other bases

Not Official
DUGONG OIL

An Oil obtained in Australia from *Halicore Dugong*, Daub, by boiling the superficial fat. A substitute for Cod-Liver Oil, recommended at one time (*P J* (3) in 3, 100) as not being disagreeable in taste, but it does not possess this character now.

Not Official
DULCAMARA

The dried young Branches of *Solanum Dulcamara* (Bittersweet), from indigenous plants which have shed their leaves.

Fresh ones have been found (*C D* '02, in 313, *Y B P* '02, 491) to contain two alkaloids, Solanidine and Solanidine, a glucoside (Solanein), and a bitter principle (Dulcamarin) of a glucosidal nature yielding on hydrolysis Dulcamaretin and Glucose.

Medicinal Properties—Alterative and sedative. Used in cutaneous eruptions, chiefly of a scaly nature, as psoriasis and pityriasis, a decoction being applied externally, at the same time that it is used internally.

An alkaloid *Solanine* obtained from *Solanum nigrum*, *S. Dulcamara* and *S. tuberosum* (Potato plant), has been recommended as an analgesic.—*L M R* '86, 496, '88, 242, *T G* '87, 56, '88, 630, *L* '87, in 1097.

Foreign Pharmacopœias—Official in Austr, Fr (*Douce-amère*), Mex, Port (*Doce-amarga*), and Span. Not in the others.

EXTRACTUM DULCAMARÆ FLUIDUM—1 fl oz equals 1 oz Dulcamara. Prepared with diluted Alcohol.—*U S P* 1890.

Dose—30 to 60 minims = 1.8 to 3.6 cc.

Foreign Pharmacopœias—Official in Mex.

A solid *Extractum Dulcamara* is official in Austr, Fl and Mex.

INFUSUM DULCAMARÆ—Dulcamara, 1, boiling Water, 10, infuse 1 hour.

Dose—1 to 2 fl oz = 28.4 to 56.8 cc.

This has been incorporated in the *B P C*.

Foreign Pharmacopœias—Official in Fl (1 in 50). Not in the others.

ELATERIUM.

ELATERIUM

Fr, ELATERION, GER, ELATERIUM, ITAL, ELATERIO, SPAN, ELATERIO

A sediment from the juice of the Fruit of *Ecballium Elaterium*.

It contains from 20 to 40 p.c. of Elaterin, to which principle the activity of the drug is due. It contains in addition a second crystallisable bitter principle, Prophetin, and the amorphous substances Ecballin or Elateric Acid, Hydroelaterin and Elateride, of which but little is at present known.

'Extractum Elaterii' was the official synonym in *B P* '85 for Elaterium.

Medicinal Properties—The most powerful hydragogue cathartic, only used in special cases. Employed in cardiac or renal dropsy and in constipation. Its administration in a debilitated state of the system or in gastro-intestinal inflammation requires very great caution on account of the depression which it produces.

Dose — $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.006 to 0.032 gramme

Prescribing Notes — *On account of the similarity in name to the active principle, care must be exercised to avoid confusion. The Pulvis Elaterii Compositus is often preferred, it is frequently given in the form of Pill with Compound Extract of Colocynthis and Henbane. To prevent it causing persistent diarrhoea, it may be given with Henbane, especially in renal diseases, in cardiac cases it should be guarded by a stimulant to prevent too much depression.*

Official Preparations — Elaterinum, Elaterin is contained in Pulvis Elaterii Compositus

Not Official — Pilula Elaterii Composita

Antidotes — The same as for Croton Oil (*q.v.* p. 460)

Foreign Pharmacopœias — Official in Mex., Elaterio, Port., Extracto de Pepinos de S. Gregorio. Not in the others.

Descriptive Notes — Two forms of Elaterium are met with in commerce, viz., English and Maltese.

The English form is very brittle, and consequently is usually met with in thin flakes or fragments $\frac{3}{4}$ inch to $\frac{1}{2}$ inch in width, the colour when fresh is green, soon becoming greyish, and when kept long or not kept dry it turns yellowish-grey. It should contain no Starch (*B.P.*), it yields up to 33 p.c. of Elaterin. The Maltese occurs in square cakes or tablets about 1 in. in diameter and rather more than $\frac{1}{2}$ in. in thickness, of a greenish grey colour, it sometimes contains Starch and Calcium Carbonate, and yields about 27 p.c. of Elaterin.

The English drug is official.

Tests — Elaterium should yield no marked effervescence on the addition of Hydrochloric Acid, indicating the absence of more than traces of Carbonates. When boiled with Water, cooled, and tested with Iodine Solution no decided blue coloration should be produced, indicating the absence of more than a trace of Starch. It is officially required to yield 50 p.c. of its weight to boiling Alcohol (90 p.c.), and when exhausted successively with Chloroform and Ether and the process repeated, it is required to yield 25 p.c. or not less than 20 p.c. of Elaterin. Good specimens of Elaterium yield from 30 to 40 p.c. of Elaterin.

Preparations

ELATERINUM ELATERIN ($C_{20}H_{28}O_5$), eq. 344.89, occurs in small hexagonal scales or tables.

It is the active principle of Elaterium.

Solubility — Insoluble in Water, sparingly in Alcohol (90 p.c.), 1 in 12 of Chloroform.

A recent figure obtained for Alcohol (90 p.c.) was 1 in 1100.

Dose — $\frac{1}{40}$ to $\frac{1}{10}$ grain = 0.0016 to 0.0065 gramme.

Foreign Pharmacopœias — Official in U.S. Not in the others.

Tests — Elaterin when heated to $190^{\circ}C$ ($374^{\circ}F$) turns yellow, and melts at $216^{\circ}C$ ($420.8^{\circ}F$). Its melting point is, however, given in the *B.P.* It should be neutral in reaction towards Litmus. A crimson colour rapidly changing to scarlet is produced on the addition of Sulphuric Acid to a solution of Elaterin in melted Phenol.

Sulphuric Acid colours it yellow, the colour gradually changing to scarlet. A crystal evaporated to dryness with a little Hydrochloric Acid leaves a residue which when washed with hot Water and subsequently treated with Sulphuric Acid produces a brownish-red colour. Mineral matter and alkaloids are the more commonly occurring impurities. The former may be detected by the residue left on the ignition of the sample with free access of air, the latter by the production of a precipitate when Tannic Acid Solution, Mercuric Chloride Test Solution, or Platonic Chloride Solution is added to a solution of the principle in Alcohol (90 p c)

PULVIS ELATERINI COMPOSITUS. COMPOUND POWDER OF ELATERIN

Elaterin, 1, Milk Sugar, 39

Dose —1 to 4 grains = 0.06 to 0.24 gramme

Foreign Pharmacopœias —Official in U.S. (Tituratio Elaterini), Elaterin, 1, Milk Sugar, 9. Not in the others

Not Official

PILULA ELATERII COMPOSITA —Elaterium, $\frac{1}{2}$ grain, Compound Extract of Colocynthis, 2 grains, Calomel, $1\frac{1}{2}$ grains, Capsicum, $\frac{1}{2}$ grain, Syrup of Glucose q s —*St. Bartholomew's*

Dose —1 or 2 pills.

Not Official

ELEMI.

A concrete, resinous exudation, the botanical source of which is undetermined, but is sometimes referred to *Canarium commune*, L. It has lately been attributed to *Canarium Luzonicum*, Miq.

It is imported from Manila.

When of good quality it is pale yellow of the consistence of stiff Honey and has a Terrene odour.

Brazilian and Yucatan Elemis are official in some of the Foreign Pharmacopœias, but are derived from other species of the same natural order *Burseraceæ*. They are usually more discoloured and harder, but have a similar odour.

Solubility —The greater part is soluble in Alcohol (90 p c), wholly soluble in Ether.

Medicinal Properties —The ointment is stimulant to indolent ulcers.

Foreign Pharmacopœias —Official in Austr., Belg., Dutch, Fr., Mex. (Goma de Liriodendron), Port., Span. and Swiss. Not in the others.

UNGUENTUM ELEMII —Elemi, 1, Spermaceti Ointment, 4, melt, strain, and stir till cold —*B.P.C.* 1885 (1 in 5)

This has been incorporated in the *B.P.C.*, using Unguentum Simplex.

Foreign Pharmacopœias —Official in Mex., Span. and Swiss, 1 of Elemi and 1 of Turpentine in 4 of Ointment, Dutch, 3 of Elemi, 2 of Turpentine in 10 of Ointment, Port., 2 of Elemi and 1 of Turpentine in 10. Not in the others.

Not Official

EMBELIA

The Fruit (including the dried Fruit and the Seeds) of *Embelia Ribes*, Burm. f., and of *Embelia robusta*, Roxb., are official in the Ind. and Col. Add. for India and the Eastern Colonies.

The powdered Seeds are used in India for tapeworm — *L* '87, n 199

Dose — 60 to 240 grains = 4 to 16 grammes

ACIDUM EMBELICUM — Obtained from the Seeds Insoluble in Water
It forms salts with Ammonium, Potassium, and Sodium

AMMONII EMBELAS — A tasteless crystalline salt, in red needles

Dose — 3 to 6 grains = 0.2 to 0.4 gramme, in Honey or Simple Syrup

Not Official

EMBLICA

The fruit of *Phyllanthus Emblica* L., (Emblie Myrobalan) has been used in Hindu medicine for a long time, as a diuretic and laxative. The fresh fruits preserved in Syrup are imported into this country

Not Official

EPHEDRINE HYDROCHLORIDE

The Hydrochloride of an alkaloid obtained from *Ephedra vulgaris*, L., or *E. Helvetica*, C. A. Mey

Has been recommended as a mydriatic in the form of a 5 p c solution —
B M J E '98, n 92

The addition to it of 1 p c of Homatropine Hydrochloride enhances its action, and the mixture, which is supplied under the name 'Mydrine' is a white powder readily soluble in Water, a 10 p c aqueous solution dilates the pupil moderately within a few minutes, without affecting the accommodation, and its effects pass away in two to four hours. It is useful in diagnostic examinations —
L '98, n 24, *T G* '98, 757

ERGOTA.

ERGOT

FR, ERGOT DE SEIGLE, GER, MUTTERKORN, ITAL, SEGALA CORNUTA,
SPAN, CORNEZUELO DE CENTENO

The sclerotium of *Claviceps purpurea*, Tulasne, in the ovary of *Secale cereale*, L.*

The drug should be stored whole, should be well dried, and kept in airtight vessels and perfectly dry. The *USP* says that after being kept for one year it is unfit for use.

The two principal varieties of Ergot are Spanish and Russian. They contain respectively about 0.2 and 0.25 p c of Ergotinine, and although the former contains somewhat less Ergotinine than the latter, it is usually considered the best. Ergot yields its virtues to Water and to Alcohol.

Ergot contains, in addition to the crystalline alkaloid Ergotinine discovered by Tanret, a second alkaloid Ergotoxine discovered by Barger and Carr of the Wellcome Physiological Research Laboratories, and described by them at the British Association meeting at

* Ergot is common on grasses, and if it grows in the pastures where cattle feed, it is said to occasion dry gangrene, causing them to lose their hoofs and horns.

York in 1906 (*C N* '06, 89, *B M J* '06, n 1792) The latter alkaloid, although itself amorphous, forms a number of crystalline salts. It is claimed to be the most important if not the one essential active principle, whilst the pure crystalline Ergotinine is almost if not quite physiologically inactive. Tanret (*J Pharm Chemie*, '06 [vi], 24, 397-403) *JCS Abs*, '06, i 979, *YBP* '07, 62) takes exception to the application of the name Ergotoxine to the amorphous body accompanying crystalline Ergotinine, which he discovered and named amorphous Ergotinine, and also strongly controverts the statement that crystalline Ergotinine is almost physiologically inactive, alleging in support the constant therapeutic employment of the base. Barger and Corn point out (*JCS Trans* '07, 340) that Tanret himself attributed the variation in the specific rotation of amorphous Ergotinine to varying amounts of crystallised Ergotinine contained in it, and that he therefore had not prepared the pure alkaloid. As to the physiological activity, they refer the reader to the experiments of H. H. Dale, published in the *Ann Physiol* '06, 34, 163. According to Barger and Corn (*JCS Trans*, '07, 339), Acetic Anhydride converts the crystalline into the amorphous alkaloid, the change being brought about by the removal of a molecule of Water and not by the introduction of an Acetyl group, and support the suggestion of Kraft's that the amorphous is the Hydrate of the crystalline alkaloid, and this theory is regarded as definitely established by their analysis. Hydroergotinine, discovered independently by Kraft (*JCS Abs* '06, i 979), is considered (*JCS Trans* '07, 341) to be undoubtedly identical with Ergotoxine. Cornutine does not occur as such in Ergot, but is an artificial decomposition product of Ergotinine. The Picrosclerotine of Dragendorff and the Socalme of Jacobi are regarded as other names for Tanret's alkaloid.

Ergot also contains Ergotinic Acid (the Sclerotic Acid of Dragendorff and the Ergotic Acid of Wenzell), Sphacelmic Acid or Sphacelotoxin, Ergosterol, Trehalose, and Mannite. Colouring matters, *eg*, Sclererythrin, Scleroidin, Sclerovanthin and its anhydride Sclerocrystallin, Picrosclerotine and Fusco-sclerotic Acid, are present, and about 33 per cent. fixed oil, which can be extracted with Ether, Petroleum Ether, and to a great extent by hydraulic pressure.

Medicinal Properties.—Ergotic, used in obstetric practice to contract the uterus, assist expulsion of placenta, and prevent or stop post-partum hæmorrhage. Employed in uterine hæmorrhage from other causes, such as fibroid tumour, and in subinvolution of the uterus, also, but with doubtful success, in hæmoptysis, hæmatemesis, hæmaturia, and epistaxis. Efficacious in flatulent dilatation of stomach, acute myelitis and in hæmorrhages of hæmorrhagic origin, in night sweats of phthisis. In hæmorrhage of the rectum gives most relief. In action in critical cases. Injections into sphincter are valuable in prolapsus ani. After elaborate investigation Robert recommends finely-powdered Ergot for certainty of

of this value have been increased by its employment in preventing abdominal surgery. 30 minims of a

specialty-prepared Ergot, in which from $\frac{1}{80}$ th to $\frac{1}{30}$ th grain of Strychnine or $\frac{1}{30}$ th grain Sparteine is dissolved, are injected three times daily, for two or three days before operation. In chorea 1 to $\frac{1}{2}$ drm of the liquid extract with 2 minims of Liquor Strychninae have been given thrice daily (*B M J* '05, i 354), or 1 drachm liquid extract and 3 minims of Liquor Arsenicalis — *B M J* '05, i 354.

It is this drug which has given the best results (*L* '05, i 851), and which seems to solve the difficulty of the treatment of surgical shock. 2 grammes of a specially prepared Ergot diluted with 20 c c of normal saline solution, 5 to 10 c c injected at a time. Its great advantage over Adrenalin is that its action is more prolonged, one dose being sufficient to keep up the blood pressure for some time.

The best remedy for intestinal hæmorrhage — *T G* '07, 324.

In hiccough (*L* '85, ii 276), in periodic neuralgia (*T G* '94, 343), in diabetes insipidus, 30 minim doses of the Liquid Extract every three hours — *L M R* '80, 231, 446, '81, 12.

In chorea 1 dram doses of the Liquid Extract given every four hours — *B M J* '03, ii 133.

Ammoniated Tincture stated to be an active preparation, and to have proved useful in obstinate cases of uterine hæmorrhage when other Ergot preparations have failed — *C D* '01, i 324, 663.

Dose — 20 to 60 grains = 1.3 to 4 grammes.

Prescribing Notes — *The unpleasant taste of the preparations of Ergot is improved by Tincture of Orange and Chloroform Water, or better by Tincture of Orange and Cinnamon Water. The Infusion and Hypodermic Injection should be made fresh as required. When the extract is ordered in pills, Powdered Liquorice Root added q s makes a good pill.*

The prescriber has three fluid extracts to choose from: (1) B P which is exhausted by cold Water, (2) U S P by diluted Alcohol mixed with Acetic Acid, (3) Liquor Ergotæ Ammoniatæ (not official) by Ammoniated diluted Alcohol. The official Tinctura Ergotæ Ammoniatæ is similar to the last, but much weaker.

It is often desired to give Iron with Ergot, which produces an unsightly ink-like mixture and a precipitate. This can be avoided by adding 6 grains of Citric Acid to 1 fl. drm of Tincture of Perchloride of Iron.

Incompatibles — Astringents, metallic salts.

Official Preparations — Extractum Ergotæ, Extractum Ergotæ Liquidum, Infusum Ergotæ, Tinctura Ergotæ Ammoniatæ. Injectio Ergotæ Hypodermica is made with Extractum Ergotæ.

Not Official — Discs of Ergotin, Extractum Ergotæ, Extractum Fungi Secalis Fluidum, Extractum Secalis Cornuti, Extractum Secalis Cornuti-Cornutino, Sphacelinicum, Fluidextractum Ergotæ, Liquor Ergotæ Ammoniatæ, Mistura Ergotæ, Mistura Ergotæ Ammoniatæ, Mistura Ergotæ et Ferri, Pilula Ergotini, Tinctura Ergotæ, Vinum Ergotæ, Acidum Scleroticum, Cornutine Citrate, Ergotin (various), Ergotinine, Ergot Aseptic, Ergotoxine, Ergotoxine Hydrochloride, Ergotoxine Phosphate.

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Ger, Hung, Jap, Norw, Russ, Swed and Swiss (Secale Cornutum), Fr (Ergot de Seigle), Ital (Segala Cornuta), Mex (Guernecillo de Centeno), Port (Cravagem de Centeno), Span (Cornezuelo de Centeno), U S (Ergota).

The Brussels Conference agreed that Ergot should be not more than one year old, and kept whole.

Descriptive Notes — Ergot is the compact horny mycelium or spawn of the small fungus, *Claviceps purpurea*, Tulasne, and is developed on, and takes the place of, the growing ovary of the rye plant. The fungus itself resembles a minute mushroom in shape, without gills, but with cavities containing ascospores, in its cap or head. It cannot be developed from the Ergot of commerce, the vitality of which is destroyed by drying, but can readily be grown on damp sand in

spring from the mature but undried Ergot. As met with in commerce Ergot may vary in size according to the variety, but averages about $\frac{1}{2}$ to $\frac{3}{4}$ of an inch (12 to 19 mm) and 1 to 2 lines in diameter. The official description limits it to $\frac{1}{3}$ to $1\frac{1}{2}$ inch (1 to 4 cm) in length, but does not give the diameter, which varies from $\frac{1}{2}$ to 4 lines. There are three principal varieties in commerce, viz., Spanish, German or Austrian, and Russian, and occasionally a variety from the Canary Islands. The Spanish is the largest and most highly priced, the German comes next in size, and then the Russian, the Canary kind containing a larger number of small specimens than the Russian. English Ergot is not known in retail commerce, it is probably mixed with the foreign drug in this country, being separated from the cereal by millers. The activity of Ergot appears to depend more upon the method of preservation than upon the particular variety employed.

It should be hard and dry. It is longitudinally furrowed on each side, violet-black externally, and pinkish-white within, with a short fracture often irregularly cracked. Specimens that are flexible have a mouldy odour, and are much cracked, usually due to exposure to damp before drying, or are infested with powder-like mites, should be rejected. If dried over lime or in a current of warm air, and kept in stoppered bottles from which air is excluded by Vaseline around the stoppers, it will keep good for some months, but should only be powdered when required for use. It should not be kept longer than a year (*P G* and *U S P*), and should not give off an ammoniacal or rancid odour when 10 parts of boiling Water are poured over it (*P G*).

Tests.—Ergot possesses a peculiar and disagreeable odour, and if it be reduced to powder and the powder is moistened with Potassium Hydroxide Solution, this odour is intensified. Good Ergot contains 3 to 5 p.c. of ash when ignited with free access of air. Specimens examined in the author's laboratory left from 2.15 to 2.96 p.c., with an average of 2.75 of ash on incineration. The cold Water extract varied from 11.04 to 13.4 p.c., with an average of 12.3 p.c.

Preparations.

EXTRACTUM ERGOTÆ.—EXTRACT OF ERGOT *B P Syn*—**ERGOTIN**

100 of Ergot exhausted by percolating with Alcohol (60 p.c.) Evaporate percolate to 25 and mix it with an equal quantity of Distilled Water, filter, add 4.7 of diluted Hydrochloric Acid, and after 24 hours filter, wash the residue in the filter until free from acid, add 2 of Sodium Carbonate to the filtrate mixed with the washings and evaporate the whole to soft extract.

Dose—2 to 8 grams *U S P* to 0.52 gramme

The corresponding preparation as to *U S P* '85 was prepared from Liquid Extract of Ergot and Rectified Spirit.

Extractum Ergotæ (U S P) is practically the same as *B P*, except that Citric Acid is added towards the end of the process. It is an alcoholic extract of Ergot treated with diluted Hydrochloric Acid to precipitate Sclerotin, the characteristic colouring matter of Ergot, neutralised with Monohydrated

Sodium Carbonate and evaporated as directed in *U S P*. One would gather from the note in the *B P C* that *U S P* employed less than half the *B P* quantity of Sodium Carbonate, but this is not so, Sodium Carbonate should read Mono hydrated Sodium Carbonate, then the quantities are nearly the same

Extractum Secalis Cornuti (*Ger*)—2 of Ergot macerated in 4 of Water, twice, the liquors mixed and evaporated to 1, mixed with 1 of Alcohol (90 p c), filtered after 3 days, and evaporated to a thick extract

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex (*Ergotina de Bongean*), Norw, Port, Russ, Span, Swed, Swiss and U S

The *Brussels Conference* agreed to prepare a watery extract and make up with Alcohol (60 p c)

EXTRACTUM ERGOTÆ LIQUIDUM LIQUID EXTRACT OF ERGOT *N O Syn*—**EXTRACTUM SECALIS CORNUTI FLUIDUM**

Macerate 20 of crushed Ergot in 100 of Distilled Water for 12 hours, strain, repeat the maceration with a further 50 of Distilled Water and strain, press the marc, strain the fluid, mix it with the other fluid portions and evaporate to 14, when cold add $7\frac{1}{2}$ of Alcohol (90 p c) and after one hour filter (In practice it is better to allow it to stand for several hours) (1 in 1)

Dose—10 to 30 minims = 0.6 to 1.8 c c

60 minims = 3.6 c c, is not infrequently prescribed

Tests—Liquid Extract of Ergot has a specific gravity of 1.015 to 1.025, contains from 13 to 16 p c w/v of total solids and about 32 p c w/v of Absolute Alcohol

Fluidextractum Ergotæ (*U S P*)—Percolate 100 of Ergot (in No. 60 powder) with 2 of Acetic Acid (*U S P*) mixed with 98 of Alcohol (49 p c), continue the percolation with Alcohol (49 p c) until exhausted, reserve the first 85, evaporate the remainder to a soft extract, dissolve this in the reserved portion and make up to 100 with Alcohol (49 p c)

Foreign Pharmacopœias—Austr (*Extractum Fungi Secalis Fluidum*) The fat is extracted from 100 of Ergot with Petroleum Ether, and after drying the marc, it is percolated with a mixture of Glycerin 5, Alcohol 20, Water 20, to produce 100, Belg, Dan, Ger, Norw, Swed and Russ, Extract with **Hydrochloric Acid** and dilute Alcohol, Fr, exhausted with Water, and Tartaric Acid, Mex (*Extracto Fluido de Curnecillo de Centeno*), with **Acetic Acid** and dilute Alcohol, Swiss and U S percolated with dilute Alcohol acidified with **Acetic Acid**, Dutch, with dilute Alcohol and Tartaric Acid, Jap, with a mixture of Alcohol 2, Water 8. Not in the others

The *Brussels Conference* agreed that the strength should be 100 p c

INFUSUM ERGOTÆ INFUSION OF ERGOT

Infuse 1 of freshly crushed Ergot wth 20 of boiling Distilled Water for 15 minutes and strain

Dose—1 to 2 fl oz = 28.4 to 56.8 c c

Used also as an injection for gleet

INJECTIO ERGOTÆ HYPODERMICA. HYPODERMIC INJECTION OF ERGOT *B P Syn*—HYPODERMIC INJECTION OF ERGOTIN

Extract of Ergot, 100 grains, Phenol, 3 grains, Distilled Water, 220 minims, or a sufficient quantity. Mix the Phenol with the Distilled Water, boil for a few minutes, cool, add the Extract of

Ergot, and, if necessary, sufficient recently boiled and cooled Distilled Water to produce 330 minims of the Injection (1 in 3)

The above is the official wording, but it is not clear why the Water should be boiled *after* the addition of the Phenol. It would be better to dissolve both the Ergot and the Phenol in the previously Sterilised Water.

Dose, by *subcutaneous injection*—3 to 10 minims = 0.18 to 0.6 c.c.

This injection should be recently prepared. 33 minims = 1 grain of Extract of Ergot.

Foreign Pharmacopœias—Official in Port (Solutio de Ergotino cum Glycerino), Ergotin 1, Glycerin 4, Water 5, all by weight, Span (Inyección Hipodérmica de Ergotina), Ergotin 1 gramme, Glycerin 2 grammes, Water *q s* to 10 c.c. Mex and Span have Injection Ergotinine.

TINCTURA ERGOTÆ AMMONIATA AMMONIATED TINCTURE OF ERGOT

Ergot, in No. 20 powder, 5, Solution of Ammonia, 2, Alcohol (60 p.c.), *q s* to yield 20 (1 in 4)

Dose—30 to 60 minims = 1.8 to 3.6 c.c.

Foreign Pharmacopœias—A simple tincture is official in Dutch, Mex and Port, 1 in 5, all by weight. Not in the others, U.S. (Vinum Ergotæ), 1 in 5.

Tests—Ammoniated Tincture of Ergot has a specific gravity of 0.930 to 0.938, contains from 3 to 5 p.c. of total solids and about 52 p.c. w/v of Absolute Alcohol.

Not Official.

DISCS OF ERGOTIN $\frac{1}{2}$ grain = 0.02 gramme, and $\frac{1}{4}$ grain = 0.016 gramme are prepared for hypodermic use.

PILULA ERGOTINI—Ergotin 2 grains, Liquorice Powder 3 grains.

LIQUOR ERGOTÆ AMMONIATUS—A liquid Extract of Ergot (1 in 1), prepared with ammoniated diluted Alcohol.

Dose—10 to 60 minims = 0.6 to 3.6 c.c.

B.P.C. gives the following formula, under the title **Extractum Ergotæ Ammoniatum Liquidum**, but adds that although four times the strength of the official tincture, it is not very active—100 of Ergot in No. 20 powder is percolated with a mixture of Solution of Ammonia 10, and Alcohol (60 p.c.) 70, continue the percolation with Alcohol (60 p.c.) until exhausted, reserve the first 85, evaporate the remainder to 15 and mix.

MISTURA ERGOTÆ—Liquid Extract of Ergot, 30 minims, Diluted Sulphuric Acid 10 minims, Chloroform Water, to 1 fl. oz.—*St. Thomas's*.

This has been incorporated in the *B.P.C.*

Liquid Extract of Ergot, 80 minims, Syrup of Ginger, 30 minims, Infusion of Orange Peel, to $\frac{1}{2}$ fl. oz.—*London*.

MISTURA ERGOTÆ AMMONIATA—Liquid Extract of Ergot, 20 minims, Ammonium Carbonate, 3 grains, Emulsion of Chloroform, 15 minims; Camphor Water, to 1 fl. oz.—*University*.

Liquid Extract of Ergot, $\frac{1}{2}$, Ammonium Carbonate, $\frac{1}{2}$, Emulsion of Chloroform $\frac{1}{2}$, Camphor Water, *q s* to produce 100—*B.P.C. Supplement*.

MISTURA ERGOTÆ ET FERRI—Liquid Extract of Ergot, 30 minims, Solution of Ferric Chloride, 15 minims; Citric Acid, 5 grains, Chloroform Water, to 1 fl. oz.—*Guy's*.

TINCTURA ERGOTÆ—Ergot, 5, Proof Spirit, 20—*B.P. 1885*.

This has been incorporated in the *B.P.C.*

VINUM ERGOTÆ—Fluid Extract of Ergot, 20, Alcohol (95 p c), 5, White Wine, 75 —*U S P*

This has been incorporated in the *B P C*, using Detannated Sherry

ACIDUM SCLEROTICUM—A weak acid principle obtained from Ergot by Dragendorff. It is used hypodermically $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.021 to 0.05 gramme, dissolved in Distilled Water or Thymol Water

CORNUTINE CITRATE—A soluble salt of an alkaloid which is stated by Kobert to be the active principle of Ergot. A brown powder, which is used in obstetric practice

Dose— $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.0054 to 0.01 gramme, or subcutaneously $\frac{1}{32}$ to $\frac{1}{8}$ grain = 0.002 to 0.008 gramme

Given in $\frac{1}{100}$ grain hypodermically to contract uterus in a case of eclampsia
—*L* '99, 1 1430

A soluble Cornutine Hydrochloride has also been prepared

ERGOTININE—An alkaloid obtained from Ergot. Long white crystals which have a tendency to darken on exposure to light and air

Solubility—According to Barger and Carr 1 part by weight of Ergotinine dissolves in 312 parts by weight of Absolute Ethyl Alcohol at 10° C (50° F) and in 292 parts by weight at 18° C (64.4° F). It is soluble 1 in 1020 parts by weight of Absolute Ether, 1 in 91 parts by weight of Ethyl Acetate, 1 in 26 parts by weight of Acetone, 1 in 77 parts by weight of boiling Benzene, 1 in 52 parts by weight of boiling Ethyl Alcohol, and 1 in 56 parts by weight of boiling Methyl Alcohol. It is stated by these authors to be extremely soluble in cold Chloroform, moderately so in Amyl Alcohol or Xylene, and insoluble in Petroleum Ether

These figures also appear in the *B P C*, but the abstractor has failed to note that all the fluids are by weight, and not by measure, and '1 in 91 parts by weight of Ethyl Alcohol' has been incorrectly copied as '1 in 91 by volume of Ethyl Acetate'

Foreign Pharmacopœias—Official in Fr (Ergotinine Cristallisée), Span and Mex

Tests—Ergotinine melts according to Barger and Carr at 229° C (444.2° F). Its solution in Ethyl Alcohol is strongly dextrogyrate, the rotation of a saturated solution in this solvent at 10° C (50° F) being + 338°, but the rotation is effected by prolonged boiling

Ergotinine is precipitated by the usual alkaloidal reagents such as Potassio-mercuric Iodide Solution, Iodo potassum Iodide Solution, Gold Chloride Solution, Platinum Chloride Solution, Bromine Water, Tannic Acid Solution and Picric Acid Solution. The addition of concentrated Sulphuric Acid to a solution of Ergotinine in Ether or in Ethyl Acetate produces a transitory orange coloration changing to blue. When the alkaloid is dissolved in concentrated Sulphuric Acid and a little anhydrous Ferric Chloride is added, a pale yellow coloration passing through orange, crimson and green to a permanent dark blue is produced

A soluble Ergotinine Citrate has also been prepared

ERGOTOXINE—This alkaloid was discovered by Barger and Carr, and is described by them as a light white powder

Solubility—It is more soluble in organic solvents than Ergotinine, notably in cold Alcohol. It is also soluble in Sodium Hydroxide solution. It is but slightly soluble in Ether

Tests—Ergotoxine, according to the above-named authors, begins to soften about 155° C (311° F) and gradually melts at 162° to 164° C (323.6° to 327.2° F). The rotation of an alcoholic solution varies with the method of preparation. Ergotoxine is precipitated by the usual alkaloidal reagents, *e.g.*, Potassio-mercuric Iodide Solution, Iodo potassum Iodide Solution, Auric Chloride Solution, Platinum Chloride Solution, Picric Acid Solution, Phosphomolybdic Acid, Bromine Water and Tannic Acid Solution. The addition of Concentrated Sulphuric Acid to a solution of Ergotoxine in Ethyl Acetate or Ether gives rise to a transitory orange coloration, changing to blue. The addition of anhydrous

Ferrie Chloride to a trace of the alkaloid dissolved in concentrated Sulphuric Acid gives a pale yellow coloration changing through orange, crimson and green to a permanent dark blue

ERGOTOXINE HYDROCHLORIDE—This salt was prepared by Barger and Carr, and is described by them as forming minute diamond shaped plates, and very thin and very long square ended needles. It melts at 205°C (401°F). It is considered very unstable and therefore very difficult to purify.

ERGOTOXINE PHOSPHATE—This salt is described as the most easily purified of the Ergotoxine salts, it is stated to crystallise in groups of needles, and when pure in isolated needles.

Solubility—Barger and Carr state that 1 part of Ergotoxine Phosphate dissolves in 313 parts by weight of cold, and in 14 parts by weight of boiling Alcohol (90 p c).

Tests—Ergotoxine Phosphate melts according to the above-named authors at 186° to 187°C (366.8° to 368.6°F). A 1 p c solution of the salt in cold Distilled Water forms a typical colloidal solution. If an equal volume of Normal Volumetric Hydrochloric, Oxalic or Acetic Acid Solutions are added to the solution, the degree of contraction in the order named, the Hydrochloric Acid forms a thick jelly, so that the tube can be inverted without the contents escaping, whilst in the case of the Acetic Acid, the solution remains fluid.

A normal and an acid Ergotoxine Oxalate have been prepared by Barger and Carr, the former described by them as elongated, a melting point of 179°C (354.2°F), the latter as minute prisms possessing a similar melting point.

ERGOTIN—This is a synonym for *B.P.* Extract of Ergot, there are also the following commercial varieties—

Ergotinum Bonjean—An aqueous reddish-brown Extract, purified by Alcohol. 1 part Extract = 5 to 6 parts Ergot.

Dose— $\frac{1}{2}$ to $4\frac{1}{2}$ grains = 0.1 to 0.8 gramme.

Ergotinum Bonjean Depuratum pro Injectione—A purified liquid for injection, $\frac{1}{2}$ parts = 1 part Ergotin Bonjean.

Dose— $\frac{1}{2}$ to $4\frac{1}{2}$ grains = 0.1 to 0.8 gramme.

Ergotin Bombelon Fluidum (Cornutinae Ergotae)—A brownish-black liquid.

Dose—80 minims = 1.8 c.c. per os, $8\frac{1}{2}$ to 8 minims = 0.2 to 0.5 c.c. subcutaneously.

Ergotin Bombelon Spissum—Soft Extract. Administered internally in Pul. form or in Solution. Ergot Bombelon Spissum, 10 grammes (or 154 grains), Aqua Laurocerasi, 7.5 grammes (or 2 fl. drms), Alcohol (90 p c), 2.5 grammes (or 42 minims), 4 to 15 drops.

Ergotinum Denzel Fluidum—A purified Extract.

Dose—3 to 10 grains = 0.2 to 0.65 gramme.

Ergotinum Kohlmann Fluidum—Brownish black fluid, miscible with Water.

Daily Dose—60 to 75 grains = 4 to 5 grammes.

Ergotinum Purum Dialysatum Wernich Spissum—A dialysed aqueous Extract of Ergot, purified by treatment with Ether and Alcohol Soluble in Water.

Dose—10 to 30 grains = 0.65 to 2 grammes.

Ergotinum Purum Dialysatum Wernich Fluidum—2 parts = 1 part of the above preparation.

Dose—10 to 60 grains = 0.65 to 4 grammes.

Ergotinum Purum Dialysatum Wernich Siccum.

Dose —22 grains = 1.4 grammes

Ergotinum Purum Siccum Wiggers—A reddish brown powder, soluble in Water

Dose — $\frac{1}{2}$ to $1\frac{1}{2}$ grain = 0.02 to 0.1 grammes

Ergotin Yvon—A brownish black fluid, prepared from fat free Ergot by exhaustion with dilute Tartaric Acid solution

Dose —10 to 20 drops internally per os, 1 c c = 16 minims hypodermically

EXTRACTUM SECALIS CORNUTI CORNUTINO-SPHACELINICUM (KOBERT)—An Extract which combines the action of Cornutine and Sphacelinic Acid, an alkaloid and a resinous body, obtained by Kobert from Ergot. It is prepared by exhausting Ergot with strong Alcohol, and evaporating the liquid to an Extract, the fatty Oil being removed by Ether

ERGOT ASEPTIC—A sterilised concentrated preparation prepared from physiologically standardised Ergot, put up in bulbs containing 1 c c representing 2 grammes or 30 grains Ergot

Not Official

ERIGERONTIS CANADENSIS OLEUM

OIL OF CANADIAN FLEABANE

A colourless, or pale yellow, mobile liquid, distilled from the fresh flowering Herb *Erigeron Canadense*, L., which grows abundantly in American Mint fields

It has a tendency to darken in colour and to become viscid on exposure to air, and rapidly becomes resinified. It should be kept in well stoppered glass bottles of a dark amber tint, in a cool atmosphere and protected as far as possible from the light

It consists almost entirely of Dextro limonene

Medicinal Properties—Diuretic, tonic, and astringent. Chiefly employed for arresting internal hemorrhage

Dose —5 to 10 minims = 0.3 to 0.6 c c every two or three hours

Foreign Pharmacopœias—Official in U.S. Not in the others

Tests—Erigeron Oil has a specific gravity of 0.850 to 0.870, an optical rotation of not below +45°, and the greater part of the oil distils about 175° C (347° F). It should be soluble in an equal volume of Alcohol (94.9 p c)

Not Official

ERYTHROL TETRANITRATE

TETRANITRIN

A colourless, crystalline solid melting at 61° C (141.8° F) prepared from Erythrol (a tetratomic Alcohol). When kept in a dark and moderately cool place it is fairly stable, but if exposed to warmth, and especially sunlight, it rapidly undergoes decomposition. It is liable to explode on percussion, and should be handled with great care

It is but slightly soluble in Water, but dissolves readily in Alcohol (90 p c) and in Ether

It is a vaso dilator and belongs to the group of which Glycerol Trinitrate (Nitroglycerin) may be regarded as the typical representative. Blood pressure experiments show that it has a less marked but more prolonged action than that substance.—*B M J* '95, ii 1213, '97, i 907, '98, i 18, 37, 248, ii 936

A list of cases treated with Erythrol Tetranitrate.—*B M J* '99, ii 1259
Dangers.—*M P* '99, 838

Dose — $\frac{1}{2}$ to 1 grain, in alcoholic solution or in the form of tablets
Tablets are made containing $\frac{1}{2}$, $\frac{1}{4}$, and 1 grain

Not Official.

ERYTHROPHLÆUM

CASCA BARK SASSY BARK

The Bark of the *Erythrophlæum guineense*, Don Introduced as a cardiac tonic in 1877

An Ordeal Bark used in West Africa It yields an alkaloid **Erythrophlæine**, the Hydrochloride of which is soluble in Water

B.P.C. Formulary 1894 had a Tincture (1 in 10), dose 5 to 10 minims = 0.3 to 0.6 c.c., but it was omitted from the 1901 edition, it is now re-introduced in *B.P.C.* 1907

ETHYL NITRITIS LIQUOR.

See under SPIRITUS ÆTHERIS NITROSI

EUCAINE. See COCAINE, p. 413**EUCALYPTI GUMMI.**

EUCALYPTUS GUM

A ruby-coloured exudation, or so-called Red Gum, from the bark of *Eucalyptus rostrata* and some other species of *Eucalyptus* Imported from Australia

Under the name of Gummi Rubrum, this has been 'Not Official' in the *Companion* since 1871

Medicinal Properties.—Astringent, principally used in diarrhoea, dysentery, and relaxed throat

This Gum adheres with great pertinacity to the mucous surfaces, and it is probably on this account that its astringency is more effective than that of Catechu Kino, etc. although it contains less astringent matter

The **Fluid Extract** is an excellent styptic, injected into the nostril, at once stops bleeding of the nose, a tablespoonful in a pint of Water forms an astringent injection for the vagina or rectum, it also forms an astringent lotion for the eyes

Dose—2 to 5 grains = 0.13 to 0.32 gramme

Prescribing Notes—Given in the form of cachets or pills
in the form of Syringes The Tincture mixes with Water and

Official Preparation—Trochiscus Eucalypti Gummi

Not Official—Extractum Gummi Rubri Liquidum, Extractum Eucalypti
 Suppositoria Gummi R Rubri, Syrupus
 Syrupus Eucalyptus Gummi Rubri,
 Gummi, Trochiscus Eucalypti Compositus, Trochiscus
 Gummi Rubri (*Squire*)

Descriptive Notes—Although one variety of *Eucalyptus* Kino is known as Red gum in commerce it is incorrectly called gum in the *B.P.* and indeed it is often called in retail commerce by the more appropriate name of *Eucalyptus* Kino The *B.P.* states that it is obtained from the bark of *Eucalyptus rostrata*, *Scaplect*, and some

other species The one named in the *B P* is unfortunately a species which does not yield the best kind According to Mr H G Smith (*P J* (4), 23, p 101) *Eucalyptus calophylla*, R. Br., yields the best obtainable in commercial quantities, the kino of *E. rostrata* being not so astringent, and its Tincture gelatinises Eucalyptus kinos contain two tannins, one giving a green colour with Ferric Chloride, and not gelatinising, and the other giving a purple colour and gelatinising when kept (See *Proc Roy Soc N S Wales, June and Aug, 1904*) The fragments or grains are described in the *B P* as transparent and ruby red, somewhat tough, adhering to the teeth and tinging the saliva red It should be soluble to the extent of 80 to 90 p c in cold Water and almost entirely in 90 p c Alcohol The Eucalyptus kino of commerce is often blackish and opaque and consists of the natural product of the trees, but there is a preparation obtainable which is made in Australia by boiling down the fresh juice collected from incisions made in the bark of the tree This is usually distinguished under the name of 'red gum' in commerce and the *B P* characters apply to it The tincture does not gelatinise It is used especially in 'red gum' lozenges on account of its purity and ready solubility Eucalyptus Gum or Kino that is allowed to dry on the tree or is picked out of the wood is often largely insoluble in Water, owing apparently to the action of an oxydase which is only destroyed by boiling

Preparation

TROCHISCUS EUCALYPTI GUMMI—EUCALYPTUS GUM LOZENGE

1 grain of Eucalyptus Gum, in each, with Fruit basis

Not Official

EXTRACTUM GUMMI RUBRI LIQUIDUM—Red Gum, 7, Water 21 dissolve, strain, and add Alcohol (90 p c), 1

Dose—30 to 60 minims = 1 8 to 3 6 c c, in a wineglassful of Water

EXTRACTUM EUCALYPTI GUMMI LIQUIDUM—Dissolve 5 of Red Gum in 18 of Distilled Water, strain, and add 2 of Alcohol (90 p c) and sufficient Distilled Water to produce 20

This has been incorporated in the *B P C* from the *B P C Formulary* 1901

SUPPOSITORIA GUMMI RUBRI—Powdered Red Gum, 5 grains, Extract Nux Vomica, 1 grain, Cocoa nut Stearin, *q s* to make one suppository

SYRUPUS GUMMI RUBRI—Liquid Extract, 20, Sugar, 12, dissolve

Dose—30 to 60 minims = 1 8 to 3 6 c c

This has been incorporated in the *B P C* under the title **Syrupus Eucalypti Gummi**

SYRUPUS EUCALYPTI ROSTRATÆ—Red Gum of *Eucalyptus rostrata*, 800 grains, Boiling Distilled Water, 9½ oz, Refined Sugar, 16 oz, Oil of Eucalyptus, 80 minims, Mucilage of Acacia, 4 fl drms—*Pharmacy Board of Victoria (C D '06, 1 110)*

Dose—30 to 60 minims = 1 8 to 3 6 c c

This has been incorporated in the *B P C* with slight alteration of the quantities as follows—

Syrupus Eucalypti Compositus.—Eucalyptus Gum from *Eucalyptus*

Rostrata, $\frac{7}{8}$, Oil of Eucalyptus, $\frac{1}{2}$, Refined Sugar, 60, Mucilage of Gum Acacia, 2, Distilled Water, *q s* to produce 100—*B P C*

TINCTURA GUMMI RUBRI—Gum, 1, Alcohol (90 p c), 4, digest and strain Mixes with Water without becoming turbid

Dose—20 to 40 minims = 1 2 to 2 4 c c

1 part of this with 6 or 8 of Water for a gargle

Tinctura Eucalypti Gummi (*B P C*)—Eucalyptus Gum, 25, Alcohol (45 p c), *q s* to make 100

TROCHISCUS EUCALYPTI COMPOSITUS—2 grains of Potassium Chlorate, $\frac{1}{2}$ grain of Powdered Cubebs and 1 grain of Red Gum in each—*Throat*

TROCHISCUS GUMMI RUBRI (*Squre*)—Made with Rose Paste This lozenge, which has been in use for about forty years, differs in appearance and flavour from that now introduced into the *B P*

Useful for relaxed throat They have also been recommended as a preventive of sea-sickness

EUCALYPTI OLEUM.

OIL OF EUCALYPTUS

A colourless, or pale yellow, oily, limpid liquid, having a characteristic aromatic odour It should contain at least 50 p c of Eucalyptol, and but very little Phellandrene It is the volatile Oil distilled from the fresh Leaves of *Eucalyptus globulus*, and other species of *Eucalyptus*

For many years the Oil from *E amygdalina* was the most esteemed variety, and was included in *B P* '85, but it is now excluded by the tests given in *B P*, '98,

The chief constituent of the Oil is Eucalyptol (Cineol), which in good Oils amounts to from 50 to 70 p c It also contains the Terpene, Dextro-pinene, in the crude Oil various Aldehydes, principally Valeric, Butyric, and Capronic Aldehydes, and in the higher boiling point fraction a laevogyrate Ester, yielding on saponification a laevogyrate Alcohol

Solubility.—3 in 1 (or less) of Alcohol (90 p c), in all proportions of Absolute Alcohol, 1 in 38 of Alcohol (60 p c), Oil, 1 in 175)

These figures have been incorporated in the *B P C*

Medicinal Properties—It is a powerful antiseptic and deodoriser, antipyretic It is used as an inhalation in cases of pulmonary gangrene, phthisis, influenza, and coryza, and internally or by inhalation to relieve the cough in chronic bronchitis, phthisis, and asthma Mixed with Iodoform as an application to hard and soft chancres, and as urethral suppository in gonorrhœa Given internally for chronic inflammation of the bladder

The following prescription of Sir R Douglas Powell (*Edm Med Jour* 05, 465) is of great service in relieving the troublesome cough of phthisis, Eucalyptol, or Pine Oil, 3 drms, Oil of Bitter Almonds, 1 drms, Spirits of Chloroform (double strength), 1 oz Ten to 15 drops to be inhaled after the first morning coughing, in the middle of the day, and in the evening

Alarming symptoms (*L* '05, 1, 903) following the internal administration of a teaspoonful of the oil, taken plain to relieve an ordinary cold Recovery

A case of recovery by Recovery—*B M J* '06, 1 1085

With Chloroform in the treatment of actylostomiasis, also in killing tape- and thread-worms—*L* '06, 1 285.

Inhalation in whooping cough—*B M J* '86, i 430 As a disinfectant, as a throat and nose spray, and as anunction in scarlet fever—*L* '95, i 861

An infusion (60 grains in 6 oz of the leaves) twice daily in the treatment of diabetes—*B M J* '02, i 1295, ii 1884, *P J* '02, ii 118

Dose— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c c

Prescribing Notes—*Given in the form of emulsion with Mucilage of Acacia and Water, or taken on Sugar Used as an inhalation or spray May be mixed with equal parts of Olive Oil for a liniment, 1 to 3 or 4 of Olive Oil as an antiseptic ununction in scarlet fever*

Official Preparation—Unguentum Eucalypti

Not Official—Fluidextractum Eucalypti, Tinctura Eucalypti, Eucalyptus Gauze, Eucalyptus Wool and Lint, Pastille of Eucalyptus, Pastille of Eucalyptol, Nebula Eucalypti, Nebula Eucalypti et Pini, Nebula Eucalypti et Menthol et Cocainæ, Parogenum Eucalyptolis, Pessus Eucalypti, Vapor Eucalypti, Vasolimentum Eucalyptoli, Eugol, Eucalypteol, Eucalyptol, Phellandrene, Oleum Eucalypti Maculatæ var Citriodora, Eudesmol

Foreign Pharmacopœias—Official in Fr (Essence d' Eucalyptus), sp gr 0.910 to 0.930, Hung, sp gr 0.914, Jap, sp gr not given, Mex (Ácete Volátil de Eucalipto), sp gr 0.905, Norw, sp gr 0.915 to 0.925, U.S., sp gr 0.905 to 0.925 at 25° C (77° F) Not in the others The Leaves are official in Belg, Dutch, Fr, Hung, Ital, Jap, Mex, Port, Span, Swiss and U.S.

Descriptive Notes—The Eucalyptus Oil of commerce is derived chiefly from *E. globulus*, Labill in Tasmania and California, etc, and from *E. amygdalina*, Labill, *E. cneorifolia*, DC, *E. dumosa*, A. Cunn, and *E. oleosa*, F v M, in South Australia

The oil of *E. amygdalina* was that originally used in medicine in Australia. It has not the Cummin flavour and odour due to Aromadendral which characterises the last three. The oil of *E. globulus* unless rectified has an unpleasant odour. Most of the oils of commerce have been rectified to free them from irritating Alcohols and Aldehydes and colouring matter, but will, nevertheless, if kept with access of air in bottles half full, in course of time resinify and thicken. Oil of *E. globulus*, adulterated with Castor Oil, has been met with in commerce.

Tests—Eucalyptus Oil has a specific gravity, according to Baker and Smith, from 0.900 to 0.925, the official figures are 0.910 to 0.930, the *USP* gives 0.905 to 0.925 at 25° C (77° F). The Report of the Committee of Reference in Pharmacy suggests that the specific gravity should be raised to 0.910. It is difficult to gauge what this means, as the minimum specific gravity is already given as 0.910. The optical rotation in a tube of 100 mm. is officially required to be from +10° to -10°. The *BP* does not include a process for the quantitative determination of the Eucalyptol (Cineol), but contents itself with the statement that it should become semi-solid on being stirred, when cold, with a third or half its volume of Phosphoric Acid (sp gr 1.75). It must be conceded that the methods available for the quantitative determination of Eucalyptol in Eucalyptus Oils are not strictly accurate. The Phosphoric Acid method for all practical purposes is sufficiently accurate to enable the comparative values of the oils to be judged. The method has been

found most useful in arranging the several members of the genus into groups.

The Phosphoric Acid method for the determination of Eucalyptol in Eucalyptus Oils was originally discovered by Mr L R Scammell, of Adelaide, South Australia, in 1892, and was the outcome of an investigation on various samples of cheap oils then being placed on the market. For about two years the process was used by Messrs Faulding and Co for the manufacture of Eucalyptol which they shipped to England. In 1894 the process was patented by Mr Scammell as Faulding's Process in England, France, Germany, and America, as well as in the Australian Colonies. With this method available, it was possible to introduce a standardised oil containing a guaranteed quantity of Eucalyptol.

The method adopted was to prepare the Eucalyptol Phosphate in a perfectly dry powdery condition by repeatedly pressing it between fresh absorbent paper, well breaking up the cake between each pressing, until finally no moisture could be detected. The Eucalyptol Phosphate thus prepared was weighed in the tared vessel in which it was to be decomposed, and from which evaporation of the Eucalyptol was not possible. Cold Water was then added, and sufficient time allowed for the Phosphate to be perfectly decomposed without heating, usually over night. The whole was then transferred to a narrow burette, graduated in $\frac{1}{10}$ of a c.c. The aqueous portion was then separated, and this, together with the warm Water used in washing the Eucalyptol, transferred to a 100 c.c. flask. When the Eucalyptol had cooled down to the room temperature, it was measured, the remaining Water run into the flask, the Eucalyptol passed through a dry filter, and the specific gravity taken, from which the weight of the Eucalyptol was calculated. The dilute acid in the flask was then made up to the mark, and 10 c.c. titrated with Semi-normal Sodium Hydroxide Solution, using Phenolphthalein Solution as indicator, and checking the results by a Lead determination.

To obtain good results with the Phosphoric Acid method, it was found necessary to keep the temperature of the bath as low as possible, using iced Water if necessary, and to add the acid slowly by drops well incorporating it with the oil. As the compound became solid it was well broken up with the rod, and ample time given for complete crystallisation to take place. Excess of Phosphoric Acid was used over that required theoretically, assuming the richest oil to contain about 73 p.c. of Eucalyptol, the determinations were made upon 10 grammes of oil.

The Phosphoric Acid method of determination is adopted by the *U.S.P.*, and a description of the process will be found under Oil of Cajuput, p. 279. An alternative method of determination is by means of Hydrobromic Acid. Absolutely anhydrous gaseous Hydrobromic Acid is passed through a measured quantity of 10 c.c. of the oil dissolved in 40 c.c. of Petroleum Ether, maintained at a freezing temperature, until a precipitate is no longer formed. The white precipitate of Cineol Hydrobromide is transferred to a pressure filter and washed with cold Petroleum Ether. The filtrate and washings

are further treated with Hydrobromic Acid, any precipitate being separately collected and added to the bulk precipitate. The Petroleum Ether is removed from the Cineol Hydrobromide by allowing it to remain for a quarter of an hour in a vacuum. It should then be rinsed with a little Alcohol into a Cassia flask and decomposed with Water. The Cineol is brought into the graduated neck of the flask and the volume read off. The figure so obtained multiplied by 10 yields the percentage by volume present in the oil. Schimmel & Co consider the Phosphoric Acid method unreliable and useless, and give a caution against its adoption. They are also of opinion that the Hydrobromic Acid cannot lay claim to reliability either, and suggest Resorcin as suitable substance for making the determination. A measured quantity of 10 c.c. of the oil is mixed in a Cassia flask with sufficient 50 p.c. Resorcin Solution to about four-fifths fill the flask. After being thoroughly shaken for 5 minutes, the uncombined oily portions are brought into the neck of the flask by adding a further quantity of the Resorcin Solution, and the volume read off. The figure multiplied by 10 yields the percentage by volume of oily constituents other than Eucalyptol (Cineol), the latter being determined by difference. Oils very rich in Cineol require to be diluted beforehand with an equal volume of Turpentine Oil in order to prevent crystallisation of the Cineol Resorcin.

It is pointed out (*CD* '08, 1 55) that within certain limits the Phosphoric Acid method proposed by Scammell gives very fair results. It is that usually adopted for the determination of Cineol. The Resorcin process is dealt with in the same reference, the opinion expressed being that it gives very disappointing results, an oil showing a Cineol content of 65 p.c. w/v by the Phosphoric Acid method indicating 82 p.c. w/v by the Resorcin method, whilst samples of Cajeput oils showing 48 to 52 p.c. w/v by the Phosphoric Acid, indicated 80 to 84 p.c. w/v by the Resorcin method. Until further important information is forthcoming, the new process cannot be accepted as giving even approximate results.

The objections to the Resorcin process recorded *CD* '08, 1 55, have been acknowledged (*CD* '08, 1 265) by Wiegand and Lehmann of Schimmel's laboratory, who state that the error is due to the influence of the terpenes and other bodies in the oil which do not distil between 170° and 190° C (338° and 374° F.). The process originally recommended has been modified so as to permit of its use for the estimation of Cineol. 100 c.c. of the oil are distilled from a Ladenburg 3-bulb flask, in such a manner that approximately 1 drop passes over every second. The Cineol content of the principal fraction boiling between 170° and 190° C (338° and 374° F.), is then determined in the manner described above in detail. The Cineol content ascertained in the fraction is then re-calculated for the original oil, and the total content in p.c. by volume is thus obtained.

The *U.S.P.* requires the oil to contain not less than 50 p.c. w/v of Cineol. The inclusion in the *B.P.* of an assay process indicating not less than 55 p.c. of Eucalyptol has been recommended. Phellandrene, if present in the oil, may be detected by mixing the sample with

twice its volume of Glacial Acetic Acid and a saturated solution of Sodium Nitrite. If Phellandrene be present, a crystalline mass will be formed.

Preparation

UNGUENTUM EUCALYPTI. EUCALYPTUS OINTMENT

Oil of Eucalyptus (by weight), 1, Hard Paraffin, 4, Soft Paraffin,
White, 5

Now 1 in 10 instead of 1 in 5

The Leaves and Oil of *E. amygdalina* are recommended by Bosisto for making the ointment

Not Official

FLUIDEXTRACTUM EUCALYPTI (US) — Eucalyptus in No 40 powder, 100, percolate with a mixture of Alcohol (95 p c), 75, and Water, 25; reserve the first 90 and evaporate the remainder to a soft extract, dissolve this in the reserved portion, and add enough menstruum to produce 100

TINCTURA EUCALYPTI—Eucalyptus Leaves, in No 20 powder, 1, Alcohol (60 p c), to percolate 5, *B P C* The *B P C Formulary* of 1901 employed Alcohol 90 p c, the foreign Pharmacopœias use an intermediate strength *U S P* fluid extract about 70 p c Alcohol

Dose — 15 to 120 minims = 0.9 to 7.1 c c

Foreign Pharmacopœias—Official in Belg, Dutch, Fr, Hung, Ital, Mex, Port, Span and Swiss, 1 in 5, Dutch, Hung and Swiss with Alcohol (70 p c), Belg, Ital and Mex with Alcohol (80 p c) Not in the others

Eucalyptus Gauze contains about 6 p.c. of the Oil, **Eucalyptus Wool** and **Lint** 5 p.c. and 10 p.c., **Pastille of Eucalyptus** containing 1 minim of Oil is made, also **Pastille of Eucalyptol** containing $\frac{1}{2}$ minim of Eucalyptol, and both of these with Cocaine $\frac{1}{20}$ grain of the Hydrochloride, also the above with $\frac{1}{2}$ grain of Menthol in each.

NEBULA EUCALYPTI—Oil of Eucalyptus, 20 minims, Liquid Paraffin, to 1 fl. oz.—*Throat*

Oil of Eucalyptus, 1, Liquid Paraffin, *q s* to produce 20 — *B P C*

Nebula Eucalypti et Pini—Oil of Eucalyptus, 5, Oil of Pine, 7 5,
Liquid Paraffin, *q s* to produce 100—*B P C*

Nebula Eucalypti et Mentholi et Cocainæ—See p 405

PESSUS EUCALYPTI—Oil of Eucalyptus, 15 minims, Oil of Theobroma,
to 2 fl drms

VAPOR EUCALYPTI.—Oil of Eucalyptus, 20 minims, Light Magnesium Carbonate, 10 grains, Water, to 1 fl oz Mix a teaspoonful in a pint of Water at 140° F for each inhalation.—*Throat*

This has been incorporated in the *B P C*

VASOLIMENTUM EUCALYPTOLI—Eucalyptol, 20, Liquid Vasoliment, 80—*Hager*

Parogenum Eucalyptolis (Eucalyptol \overline{n} 1, 20,
Parogen, 80 — BPC

Eugol is a liquid containing Beta-naphthol, Boric Acid, Menthol, Thymol, Eucalyptol, Gaultheria, and *Hamamelis* — *B.M.J.* '98, 1 702, *L.* '98, 1 37

Eucalypteol (C₁₅H₂₆O) (chloride) —A crystalline substance, almost insoluble in Water, m.p. 170°C (320°F) and boiling at 115°C (239°F)

Dose—5 grains = 0.32 gramme, as an internal antiseptic. 80 grains in Olive Oil may be given as an enema in diarrhoea.

EUCALYPTOL (Crystallisable)—A definite chemical body ($C_{15}H_{26}O$ eq 152.98), obtained from Eucalyptus Oil by a freezing process, or by separation as

Eucalyptol Phosphate and subsequent decomposition of this salt by hot Water It is liquid at ordinary temperatures, but crystallises about 0°C (32°F) It should be kept in well closed glass bottles of a dark amber tint and protected as far as possible from the light It is identical with an oxidised compound obtained from Oil of Cajuput and a number of other essential oils, consequently the names **Cineol** and **Cajuputol** have also been applied to it

Dose —1 to 3 minims = 0.06 to 0.18 c c

Official in Fr, Belg, Ital, Port, Span, Swed, Swiss and U S

Tests —Eucalyptol has a specific gravity of 0.928 to 0.930, *U S P* 0.925 at 25°C (77°F), *Fr Codex* (1908) 0.940 at 0°C (32°F) It boils at 176° to 177°C (348.8 to 350.6°F), *Fr Codex* 176°C (348.8°F) It is optically almost inactive It is liquid at ordinary temperatures, but crystallises about 0°C (32°F) When placed in a freezing mixture and gradually mixed with an equal volume of Phosphoric Acid (1.75 sp gr) it sets to a solid white crystalline mass No diminution in volume should occur when the sample is shaken with an equal volume of Sodium Hydroxide Solution It dissolves readily in Alcohol (90 p c), forming a solution which should be neutral in reaction to Litmus paper, and which should yield no brownish or violet colour on the addition of a drop of Ferric Chloride T S

The percentage of Cineol (Eucalyptol) may be determined by the Phosphoric Acid process given under Oil of Eucalyptus

PHELLANDRENE —A levogyrate terpene, occurring in the Oil from *E amygdalina* Its presence can readily be detected by the formation of a crystalline nitrosite when the Oil is treated with Nitrous Acid

OLEUM EUCALYPTI MACULATÆ VAR CITRIODORA —A pale yellow oily liquid with a pleasant citronella like odour Sp gr 0.870 to 0.905 It contains from 84 to 90 p c Citronellal, $\text{C}_{10}\text{H}_{18}\text{O}$

EUDESMOL —A crystalline Camphor from Eucalyptus Oil

EUONYMI CORTEX.

EUONYMUS BARK

FR, FUSAIN NOIR POURPRÉ, GER, SPINDLEBAUM, ITAL, EVONIMUS, SPAN, BOMETTRO

The dried Root-bark of *Euonymus atropurpureus* Jacq

Medicinal Properties —Tonic, cathartic, and diuretic The dry extract is a powerful cholagogue and purgative, useful in chronic constipation and torpid liver

Prescribing Notes —Dried Extract in one form or another has been known for many years as **Euonymin**, usually given in the form of pills with Extract of Henbane, if prescribed alone, a little Soap, $\frac{1}{2}$ grain in a 2 or 3 grain pill, and Alcohol (90 p c) q s makes a good mass Also prescribed with Iridin, the dose of which is the same

Official Preparation —Extractum Euonymi Siccum

Not Official —Elixir Euonymini Comp, Extract Euonymi, Fluidextractum Euonymini, Liquor Euonymini, Liquor Euosmici et Pepsini, Liquor Euonymi Bismuth Pepsin cum Iridino, Pilula Euonymini et Cascara, Tinctura Euonymi

Foreign Pharmacopœias —Official in Fr and U S Not in the others

Descriptive Notes —It is probable that a part of the bark of commerce is derived from *E americanus* L, which has warty fruits and almost sessile, thick leaves The root-bark usually occurs in small curved or slightly quilled pieces $1\frac{1}{2}$ to 2 inches (37 to 50 mm)

long and $\frac{1}{2}$ to $\frac{3}{8}$ of an inch (2 to 4 mm) thick and 12 to 15 mm in width, of an ashy or brownish-grey colour externally, with scattered patches of soft cork, and occasional small transverse scars and darker lines or patches. The inner surface is pale, of a light brown colour, and the fracture is short and yellowish, with projecting silky threads, more evident when the fractured edges are gently separated. The taste is bitter, somewhat acrid, and mucilaginous. Although only the root-bark is official, both in the *BP* and the *USP*, that of the stem is also sold either separately or mixed with it. It can be distinguished by occurring in longer, thin quills, with a greenish cortical portion, a fibrous bast, and more fibrous fracture.

The bark of *Alstonia scholaris*, R. Br., has been offered for *Euonymus*, but it is twice as thick, and its transverse fracture does not show cottony threads but small granular masses of stone cells.

Preparation

EXTRACTUM EUONYMI SICCUM. DRY EXTRACT OF EUONYMUS

Exhaust *Euonymus* Bark by percolation with Alcohol (45 p c), evaporate the percolate to dryness, and to each 4 of product add 1 of Calcium Phosphate. As it is hygroscopic, it should be kept in stoppered bottles.

Dose — 1 to 2 grains = 0.06 to 0.13 gramme

Fr., a powder, U.S., an extract

Not Official

EXTRACTUM EUONYMI (U.S.) — 100 of fluid extract is evaporated to dryness, and when powdered mixed with sufficient powdered Lignum to make 25 by weight.

FLUIDEXTRACTUM EUONYMINI (U.S.) — 100 of *Euonymus* in No. 40 powder is exhausted by a mixture of Alcohol (95 p c), 80, and Water, 20, reserve the first 80 of percolate, evaporate the remainder to a soft extract, which dissolve in the reserved portion, and make up to 100.

LIQUOR EUONYMINI — *Euonymin*, 32 grains, Oil of Coriander, 2 minims, Alcohol (45 p c), 1 fl. oz. — *Bourne's Formulary*

Dose — 15 to 30 minims = 0.9 to 1.8 c c

Dry Extract of *Euonymus*, 6, Oil of Coriander, 0.75, Alcohol (45 p c), q s to produce 100 — *B.P.C.*

LIQUOR EUONYMIN ET PEPSINI — Soluble Scale Pepsin, 32 grains, Dilute Hydrochloric Acid, 80 minims, Solution of *Euonymin*, 4 fl. drms, Alcohol (45 p c), 4 fl. drms, Chloroform Water, q s to make 2 fl. oz. — *Bourne's Formulary*

LIQUOR EUONYMINI CUM PEPSINO — Tincture of *Euonymus*, 2½ fl. oz., Pepsin, in scales, 24 grains, Diluted Hydrochloric Acid, 3 fl. drms, Glycerine, 3 fl. oz., Distilled Water, to 20 fl. oz. — *A.P.H.*

This has been incorporated in the *B.P.C.* under the title **Elixir Euonymi et Pepsinæ**, the figures being 12.5, 2.0, 15, 100 respectively.

LIQUOR PEPSINI BISMUTHI ET EUONYMI CUM IRIDINO — Glycerole Pepsin (*Armour's*), 24 fl. oz., Ammonio Citrate of Bismuth, 320 grains, Tincture of *Euonymus* (*B.P.C.*), 400 minims, Iridin, 16 grains, Tincture of Cochineal q s, Simple Elixir, q s to make 20 fl. oz. — *Armour's Form.*, also *P.J.F.*

Elixir Euonymi Compositum—Tincture of Euonymus, 4, Iridin, $\frac{1}{2}$, Stronger Glycerine of Pepsine, $12\frac{1}{2}$, Bismuth and Ammonium Citrate, $8\frac{1}{2}$, Solution of Cochineal $q\ s$, Simple Elixir, $q\ s$ to produce 100—*B P C*

PILULÆ EUONYMINI ET CASCARÆ—Euonymin, 12 grains, Extract of Cascara, 36 grains, Green Extract of Hyoscyamus, 12 grains, Iridin, 12 grains, Extract of Nux Vomica, $1\frac{1}{2}$ grains Divide in 24 pills—*Pharm Form*

Pilulæ Cascaræ et Euonymini—Extract of Cascara Sagrada, $\frac{1}{2}$ grain, Euonymin, $\frac{1}{2}$ grain, Green Extract of Hyoscyamus, $\frac{1}{2}$ grain for 1 pill—*B P C*

TINCTURA EUONYMI—Euonymus Bark, in No 20 powder, 4, Alcohol (90 p c), sufficient to percolate 20—*B P C Formulary* 1901

Dose—10 to 40 minims = 0 6 to 2 4 c c

This has been incorporated in the *B P C*

Not Official

EUPATORIUM

THOROUGHWORT BONESET

The dried leaves and flowering tops of *Eupatorium perfoliatum* L. A perennial plant indigenous to the United States, and is official in *U S P*

Medicinal Properties—A bitter tonic and diaphoretic. In large doses, emetic and aperient. Has been used in bronchial catarrh, influenza, and muscular rheumatism.

FLUIDEXTRACTUM EUPATORII (U S)—A 1 in 1 fluid extract of the above prepared by percolation with Alcohol (49 p c)

Dose—20 to 60 minims = 1 2 to 3 6 c c

Not Official

EUPHORBIIUM

The concrete resinous Juice of *Euphorbia resinifera*, Beng (a native of Morocco), and other species. Official in Austr, Belg, Dan, Fr, Ger, Hung, Ital, Mex, Norw, Port, Span, Swed and Swiss. It was formerly official in London, Edinburgh and Dublin Pharmacopœias. It contains an acrid Resin. It is a powerful irritant and vesicant, and is used principally in veterinary medicine. It is noticed here because it is official in most of the Foreign Pharmacopœias. A Tincture, 1 in 5, is official in Port.

It must not be confounded with the following—

EUPHORBIA PILULIFERA—A plant growing in Queensland and tropical America. The herb is collected when in flower and carefully dried. It yields its virtues to Alcohol and to Ether.

Given in spasmodic asthma and bronchial affections, in coryza and hay fever, and in spasmodic dyspnoea of whatever origin. *L* '85, 11 '86, *T G* '85, 92, *M A* '93, 260, '94, 20, *Y B T* '94, 32.

EXTRACTUM EUPHORBIAE PILULIFERÆ—Obtained by the evaporation of the following Tincture.

Dose— $\frac{1}{2}$ to 1 grain = 0 032 to 0 065 grainme

TINCTURA EUPHORBIAE PILULIFERÆ—Euphorbia in No 20 Powder, 1, Alcohol (60 p c), to percolate, 5. *B P C Formulary* 1901, incorporated in the *B P C* under the title *Tinctura Euphorbiæ*.

Dose—10 to 80 minims = 0 6 to 8 c c, well diluted with Water.

Foreign Pharmacopœias—Not in any.

Not Official

EXALGIN

METHYLACETANILIDE

eq 148 01

Long, colourless prismatic needles, or in tabular crystals

It may be prepared by the action of Acetyl Chloride on Monomethylaniline

Solubility —1 in 50 of Water, 1 in 2 of Alcohol (90 p c), 1 in 4 of Alcohol (60 p c), 1 in 2 of Chloroform, 1 in 10 of Ether

In hot Water Exalgin is very apt to form supersaturated solutions, which when cold will not separate even when stirred or scratched, but set solid at once on the addition of a fragment of a crystal

Medicinal Properties —In small doses it acts as an analgesic without effects, giving the best results in neuralgia and toothache. It is antipyretic —*B M J* '90, 1 844, 558, '90, 1 735, *P J* (3) xix 781, 861, *T G* '89, 339, 534, 746, 797, *L* '89, 1 658, '90, 1 845, '92, 1 1174, 1175, '93, 1 785. In large doses it possesses toxic properties

Severe toxic symptoms in an asthmatic woman after taking one dose of 5 grains —*L* '95, 1 1807

Caution —by large dose (150 grains), recovery. Treatment consisted of 30 grains of Acid by nasal tube, and of injection of $\frac{1}{10}$ grain Atropine, followed by two injections of $\frac{1}{10}$ grain — 890

Dose — $\frac{1}{2}$ to 1 grain = 0.032 to 0.065 gramme, was found sufficient by Fraser, but larger doses, 4 to 8 grains = 0.26 to 0.52 gramme, have been given in France

Prescribing Notes —May be given in Mixtures, previously dissolving it in a little of the Tincture before adding the Water. A nice pill mass is made by adding Grains 15 or $\frac{1}{2}$ grain Compound Tragacanth Powder to each 3 grains of Exalgin. Syrup q s. It may also be conveniently given in cachets Compr. also prepared

Foreign Pharmacopœias —Official in Fr (Methylacetanilide), Mex and Span. Not in the others

Tests —Exalgin possesses a melting point of 101° C (213° F) and a boiling point of about 245° C (473° F). When boiled with Sodium Hydroxide solution it is decomposed with difficulty, but is completely decomposed by concentrated Hydrochloric Acid with formation of Acetic Acid and Methylaniline. When a small quantity is boiled with Hydrochloric Acid, and the cooled mixture is treated with an excess of Ammonia Solution, no violet coloration should be produced on the addition of Chlorinated Lime Solution. When boiled with a few drops of Chloroform and some Alcoholic Potassium Hydroxide Solution, no odour of Phenylisocyanide is evolved. Exalgin dissolves readily in Chloroform, and this fact enables it to be distinguished from Acetanilide and Phenacetin. When 1 gramme of the sample is treated with 2 cc of Chloroform, the Exalgin is dissolved. A chloroformic solution of Exalgin remains clear when diluted with 10 times its volume of Petroleum Ether, whereas solutions of Acetanilide and Phenacetin become turbid. 0.5 gramme should leave no weighable residue when heated with free access of air.

MISTURA METHYLACETANILIDI —Methylacetanilide, 3 grains, Syrup of Orange, 1 fl drim, Chloroform Water (*B P* '85) to 1 fl oz

FEL BOVINUM PURIFICATUM.

PURIFIED OX BILE

Evaporate 20 fl oz of fresh Ox Bile to 5 fl oz, and mix it with 10 fl oz of Alcohol (90 p c), separate the precipitate, and reduce the clear fluid to a thick extract

Solubility—Soluble in Water and in Alcohol (90 p c) In soluble in Ether

Medicinal Properties—Intestinal antiseptic and cholagogue, purgative Used where there is a deficiency of bile, it assists the emulsification of fats

Dose—5 to 15 grains = 0.32 to 1 gramme

As it is desirable that it should pass into the small intestine unchanged, the pills should be coated with Keratin solution, p 710, which protects them from the action of the gastric juice

Foreign Pharmacopœias—Official in Dutch and Jap (Fel Tauri Inspissatum), Mex (Hiel de toro) Port (Extracto de Fel de Boi), Gall 1, Alcohol 1, Animal Charcoal $\frac{1}{10}$, U S (Fel Bovis Purificatum), Ox Gall 3, concentrated to 1, Alcohol 1 Not in the others

Tests—Purified Ox Bile is soluble in Water and in Alcohol (90 p c), when dissolved in from twenty to thirty times its weight of the former liquid, and mixed with a drop of a fresh syrup prepared by dissolving one part of refined Sugar in four of Water, it yields on the addition of Sulphuric Acid cautiously added so that the precipitate at first formed is redissolved, a cherry-red colour changing through carmine and purple to violet The reaction is known as Pettenkofer's test, and is the characteristic reaction of Cholalic Acid It may also be equally well observed by treating a drop of an aqueous solution of the bile on a porcelain surface with a drop of a solution of Cane Sugar, and adding a drop of strong Sulphuric Acid Owing to a chance of the reaction being obscured by the charring of the Sugar, it has been proposed to employ Furfural or Glucose in the place of Cane Sugar Unpurified Ox Bile, if present, is revealed on the addition of Alcohol (90 p c) to an aqueous solution, if absent, no precipitate should be produced The *BP* does not state the strength of the aqueous solution to be employed nor the quantity of Alcohol (90 p c) to be added, the *USP* states that an aqueous solution of the purified Ox Gall (presumably 1 in 100) should be clear and should remain transparent upon the addition of an equal volume of Alcohol (94.9 p c)

FERRUM.

IRON

Fe, eq 55.60

FR, FER, GER, GEPULVERTES EISER, ITAL, FERRO, SPAN, HIERRO

Annealed Iron wire, having a diameter about 0.005 inch = 0.1 mm (about No 35 wire gauge), or wrought iron nails, free from Oxide

The use of Iron in medicine is of great antiquity, it is said to have been the first mineral used internally, more than 3000 years ago

Iron salts naturally divide into two groups the Ferrous or Protosalts, based upon the Oxide FeO , and the Ferric or Sesquisalts (Persalts), based upon the Oxide Fe_2O_3 Ferrous salts have a strong tendency to pass into the Ferric condition by absorption of atmospheric Oxygen, a change which takes place very rapidly in presence of oxidising agents, as Chlorine, Nitric Acid, etc

Medicinal Properties.—The Iron salts in general are hæmâtine and tonic, the Perchloride and Sulphate are also very astringent and hæmostatic, and are antiseptic. All the Iron salts are stated to be converted into Chloride by the acid of the stomach. The Astringent salts are the most powerful tonics, but as they frequently produce gastric irritation, the Neutral salts are far more generally prescribed. Of these Ferrous Carbonate in its various forms, and the Iron and Ammonium Citrate, are in the greatest demand. The Phosphate preparations are excellent hæmâtinics, and are very popular with children. Iron preparations are given after food.

The Iron and Quinine Citrate, Arsenate, and Iodide are given in special cases calling for these combinations.

Iron is useful in most forms of anæmia, and in dyspepsia, debility, chronic cachectic conditions, neuralgia, amenorrhœa and other conditions which so often depend on anæmia, also in convalescence. It is contra-indicated in apoplectic persons and generally in fevers, but has been given with benefit in erysipelas.

When constipation is a symptom, the Iron is combined with some aperient, such as Aloes and Nux Vomica or Cascara, or a mixture containing Magnesium or Sodium Sulphate may be taken separately as required.

Official Preparations.—Of metallic Iron, Ferri Sulphas, Liquor Ferri Pernitrat, Liquor Ferri Perchloridi Fortis, of Iron Wire, Syrupus Ferri, Syrupus Ferri Phosphatis, Syrupus Ferri Phosphatis cum Quinina et Vinum Ferri, of Ferrous Sulphate, Ferri Arsenas, Ferri Carbonas Saccharatus, Ferri Phosphas, Ferri Sulphas Exsiccatus, Liquor Ferri Persulphatis, Mistura Ferri Composita, of Strong Solution of Ferric Chloride, Liquor Ferri Perchloridi, Tinctura Ferri Perchloridi, of Solution of Ferric Sulphate, Ferri et Ammonia Citras, Ferri et Quinina Citras, Ferrum Tartaratum, Liquor Ferri Acetatis, of Exsiccated Ferrous Sulphate, Pilula Ferri, Pilula Aloes et Ferri, of Reduced Iron, Trochiscus Ferri Redacti, of Iron and Ammonium Citrate, Vinum Ferri Citratis.

Not Official.—Mistura Ferri Aromatica, Extractum Ferri Pomati, Iron Malate Wine, Syrupus Ferri Pomati Compositus, Tinctura Ferri Pomati.

Foreign Preparations.—in Austr, Dan, Dutch, Ger, Hung, Ital, Pruss, Russ, Siam, Spain, Sweden, Switzerland, and U.S. (see also Vinum Pulveratum), Belg (Ferri Pulvis), Fr. (Fer), Ital (Limatura de Ferro), also (Ferro Polfirizzato), Port (Ferro), Mex (Fierro), Span (Hierro), and U.S. (Ferrum).

Tests.—Iron when present in solution in the Ferric condition answers the following distinctive tests.—The addition of Ammonia Solution produces a reddish-brown flocculent precipitate, insoluble in excess of the reagent, soluble in Citric or Tartaric Acid, Potassium or Sodium Hydroxide Solution produces a similar precipitate also soluble in Citric or Tartaric Acid, Potassium Ferrocyanide Solution produces a fine blue precipitate insoluble in dilute Hydrochloric Acid, soluble in Oxalic Acid, decomposed by Potassium or Sodium Hydroxide Solution, Potassium Ferrocyanide Solution produces a brown or reddish-brown coloration but no precipitate, Ammonium Hydrosulphide Solution produces a black precipitate mixed with Sulphur, on the addition of Hydrochloric Acid the black precipitate dissolves, evolving Hydrogen Sulphide gas and leaving a white insoluble precipitate of Sulphur, Ammonium or Potassium

Thiocyanate Solution yields a blood red coloration readily destroyed by Mercuric Chloride Test-solution, also destroyed by Phosphoric Acid, Tannic Acid Solution produces a black or bluish-black coloration in dilute solutions, a black or bluish black precipitate in stronger solutions, a solution of the Ferric salt acidified with Hydrochloric Acid liberates Iodine when added to a solution of Potassium Iodide, this reaction has been utilised in the *USP* as a general method for the determination of Iron in the Ferric condition

When present in the Ferrous condition its solution yields the following reactions —Ammonia Solution produces a white flocculent precipitate, rapidly turning to a dirty green colour and ultimately to reddish brown, it is soluble in diluted mineral acid and in Citric or Tartaric Acid, rapidly becoming brown on exposure to air, Potassium or Sodium Hydroxide Solution yields a similar precipitate which behaves similarly with the reagents mentioned, Potassium Ferrocyanide Solution produces a bluish-white precipitate insoluble in dilute Hydrochloric Acid, the precipitate rapidly changes to dark blue on exposure to air, Potassium Ferricyanide Solution produces a dark blue precipitate, insoluble in dilute Hydrochloric Acid and decomposed by Potassium or Sodium Hydroxide Solution, Ammonium Hydrosulphide Solution yields a black precipitate soluble in cold diluted Hydrochloric Acid with the evolution of Hydrogen Sulphide gas, but no precipitate of Sulphur remains, Hydrogen Sulphide Solution yields no precipitate in an acid solution of a Ferrous salt, Ammonium or Potassium Thiocyanate Solution produces no reaction in solutions containing pure Ferrous salt

Preparation

VINUM FERRI IRON WINE

Iron, in wire, 1, Sherry 20 Set aside for thirty days in a closed vessel, the Iron wire being almost, but not quite, immersed in the Sherry, the vessel being frequently shaken, and the stopper occasionally removed, filter

The quantity of Iron dissolved seems to depend almost wholly upon the acidity of the Wine We found that a good dinner Sherry, containing acids equal to 0.896 p.c. of Acetic Acid, dissolved 0.14 p.c. of Iron, and had its acidity reduced to 0.09 p.c. It was treated as directed in the *BP*, and the bottle was about half full

Of such a Vinum Ferri, 3 fl. drms would represent the Iron contained in 5 minims of Tinctura Ferri Perchloridi

Commercial samples seem to lie between 0.2 and 0.3 p.c. of Iron, although occasionally samples are found much weaker

According to *PJ* (8) xxi 641, the Iron strength increases for three weeks and then diminishes Our experience does not agree with this A gallon quantity was put on and examined after the first week, and afterwards every month for four months, with the following results 0.084, 0.114, 0.157, 0.185, 0.204 p.c. of metallic Iron

N.B. —The old Vinum Ferri, made with Malaga, is much sweeter than that of the *BP*, and is sometimes ordered on that account

Dose —1 to 4 fl. drms = 3.6 to 14.2 c.c.

Prescribed for young children and delicate females with irritable stomach

Not Official

MISTURA FERRI AROMATICA—Fine Iron Wire, 2, Red Cinchona Bark, in powder, 4, Calumba, in coarse powder, 2, Cloves, bruised, 1, Compound Tincture of Cardamoms, 12, Tincture of Orange Peel, 2, Peppermint Water, 48. Macerate the first four ingredients in the last one for three days in a closed vessel, agitating occasionally, filter, and make up with Peppermint Water to 50, to this add the Tinctures, and preserve in a well-stoppered bottle—*B P* 1885

Dose—1 to 2 fl oz = 28 4 to 56 8 cc

Much valued, especially in Dublin, as a stomachic tonic and hæmatinic

This has been incorporated in the *B P C*, slightly modifying the quantities in the transposition to the decimal system

EXTRACTUM FERRI POMATI—Sour Apples, 50, convert them into a pulp and express, to the expressed liquid add Iron Wire, 1, heat the mixture on a water-bath until the evolution of gas ceases. Dilute the liquid with Water to make 50 parts, and set it aside for several days, then filter and evaporate to a thick extract. The extract should be a greenish-black, and should form a clear solution with Water

Dose—3 to 10 grains = 0 20 to 0 65 grammes

Foreign Pharmacopœias—Official in Hung (*Ext Malatis Ferri*), Austr, Dan, Norw and Swed (*Ext Pomi Ferratum*), Belg and Ger (*Ext Ferri Pomati*), Jap, Russ and Swiss (*Ext Ferri Pomatum*). Swiss is prepared by dissolving freshly precipitated Peroxide of Iron in Apple Juice, all the others are with metallic Iron and Apple Juice

SIRUPUS FERRI POMATI COMPOSITUS—Ferrated Extract of Apples, 1, Cinnamon Water, 4, Syrup of Orange Peel, 20, Simple Syrup, 24, Syrup of Rhubarb, 50, Tincture of Cinnamon, 1—*Swiss*

TINCTURA FERRI POMATI—Ferrated Extract of Apples, 1, Alcohol (90 p c), 1, Cinnamon Water, to make 10

Dose—30 to 90 minims = 1 8 to 5 4 cc

Foreign Pharmacopœias—Official in Austr, Dan, Hung, Norw and Swed, 1 and 5, Belg, Ger, Jap, Russ and Swiss, 1 and 9, Dutch (*Solutio Ferri Pomata*), and Ital (*Tinctura di Malato di Ferro*). Not in the others

IRON MALATE WINE—In Devonshi . . . Iron Wire or Nails
is digested in a bottle of Cider for a week, a . . . times a day is the
dose

FERRI ACETATIS LIQUOR.

SOLUTION OF FERRIC ACETATE

A dark brownish-red liquid possessing an odour of Acetic Acid and an acid pungent taste.

Medicinal Properties—Has a diuretic in addition to a hæmaturic and astringent action, and being compatible with Potassium Acetate, is used in some cases of Bright's disease

In the treatment of either broncho or lobar-pneumonia (*B M J* '05, 1 812), the following prescription has yielded surprisingly good results. *Liquor Ferri Perchlor*, 15 minims, *Liquor Ammon Acet*, 2 drms, *Aqua Chloroformi*, to $\frac{1}{2}$ oz, every four hours when taken alone, or every six hours when alternated with the following Strychnine mixture—*Liquor Strychninæ*, 5 minims, *Chloroform Water*, to $\frac{1}{2}$ oz

Dose—5 to 15 minims = 0 3 to 0 9 cc

Not Official—*Tinctura Ferri Acetici Ætherea*

Incompatibles—The same as given under *Tinctura Ferri Perchloridi*

Foreign Pharmacopœias—Official in Russ and Swiss, sp gr 1 087 to 1 091 Not in the others

Solubility—Miscible in all proportions with Water and Alcohol (90 p c)

Tests—Solution of Ferric Acetate has a specific gravity of 1 031 to 1 035 It is officially required to answer the tests distinctive of Ferric salts, given under strong Ferric Chloride Solution, but it should be noted that this solution will not react with Potassium Sulphocyanide Solution except in the presence of a free mineral acid (not Phosphoric), neither will it liberate Iodine from Potassium Iodide Tannic Acid Solution yields a bluish-black coloration or precipitate Ammonium, Potassium or Sodium Hydroxide Solution produces a reddish-brown precipitate, soluble in Citric or Tartaric Acid Solution When heated with Sulphuric Acid and a little Alcohol (90 p c) the characteristic odour of Ethyl-Acetate is evolved When warmed with Sulphuric Acid alone it evolves a strong acetous odour

The more generally occurring impurities are Ammonium, Arsenic, Calcium, Copper, Lead, Sodium, Potassium, Zinc, Nitrates, Sulphates, and Ferrous salts Arsenic may be detected by the modified Gutzeit's test, Copper, Lead and Zinc by Hydrogen Sulphide in either acid or alkaline solution, Ammonium by the evolution of an odour of Ammonia when the liquor is warmed with Potassium or Sodium Hydroxide Solution, Calcium by the precipitate or cloudiness produced by Ammonium Oxalate Solution, Nitrates and Sulphates after the removal of the Iron, the former by the Ferrous Sulphate ring test, the latter by Barium Chloride Solution

Not Official

TINCTURA FERRI ACETICI ÆTHEREA (*Swiss*)—Solution of Iron Acetate (sp gr 1 087 to 1 091), 8, Alcohol, 1, Acetic Ether, 1 All by weight

Dose—10 to 20 minims = 0 6 to 1 2 c c

Official in Russ, the proportions being 9, 2, and 1 respectively

Not Official

FERRI ALBUMINAS

A liquor is official in the Dutch Pharmacopœia containing 0 25 p c of Ferric Oxide, and several other formulas have been proposed, but it is more convenient to use the commercial scale preparation, which is fairly soluble in Water, and contains 5 p c of Ferric Oxide

Medicinal Properties—Hæmatinic tonic Given with success in anæmia, and specially recommended in gastric ulcer—*T G* '86, 399, *L* '94, n 1113, '95, 1 1065, *B M J E* '94, 1 28, 96, *P*, l m 87

Dose—3 to 10 grains = 0 2 to 0 65 gramme

Foreign Pharmacopœias—Official in Dan, Dutch, Ger., Jap, Russ and Swiss (*Liquor Ferri Albuminati*), Swed (*Liquor Oxydi Ferrici Albuminati*) All containing 0 4 p c of Iron

LIQUOR FERRI ALBUMINATI—Dry Egg Albumen, 4, Solution of Ferric Oxychloride, 13, Alcohol (90 p c), 12, Aromatic Elixir (*U S P*), 40,

Solution of Sodium Hydroxide (*U S P*) and Distilled Water, *q s* of each to produce 100 — *U S N F*

This has been incorporated in the *B P C*, employing 12.5 of Alcohol (90 p c) in place of 12 of Alcohol (95 p c)

FERRATIN — A brown, tasteless powder, containing 7 p c of Iron, prepared from egg Albumen and Tatarated Iron in alkaline solution. Daily dose for children, 5 to 15 grains, and for adults, 20 to 30 grains — *P*, 11 427, *A J P* '94, 500, *B M J* '95, 1 985, *B M J E* '95, 11 16, '96, 1 8, *T G* '96, 40, *L* '96, 11 1820, *B M J E* '02, 11 11

Official in Russ

Alboferin (Iron Albuminate) — An almost odourless, brown powder, soluble in Water — *B M J E* '02, 1 68

Carniferrin — A compound of Iron with Phospho-carnic Acid. A brown powder containing about 30 p c of Iron

Fersan (Iron Paranucleo-proteid) — An Iron compound, obtained from red blood corpuscles, soluble in Water — *B M J E* '00, 11 20

Dose — 10 to 30 grains = 0.65 to 2 grammes

Ferri Alginas (Alginoid Iron) — A tasteless, brown powder, containing about 10 p c of Iron. Insoluble in Water, soluble in Ammonia. Recommended in anæmia — *P J* '98, 11 199, *B M J* '02, 1 723

Claimed (*M P* '05, 11 9) that this drug has two advantages over other compounds of Iron: (1) it does not derange digestion, (2) it does not cause constipation. Alginic Acid is a nitrogenous acid obtained from seaweed. It is best given in powder or cachets.

Dose — 2 to 15 grains = 0.13 to 1 gramme

Particularly useful in chlorosis with vomiting and pain in the stomach

FERRI PEPTONAS — A brown or reddish-brown powder, having a meaty and somewhat disagreeable odour. Readily soluble in Water

Dose — 5 to 10 grains = 0.32 to 0.65 gramme

LIQUOR FERRI PEPTONATI — Peptone, dry, 4, Solution of Ferric Oxychloride, 20, Alcohol, 12, Aromatic Elixir (*U S P*), 40, Solution of Sodium Hydroxide (*U S P*) *q s*, Distilled Water, *q s* to produce 100 — *U S N F*

This has been incorporated in the *B P C*

LIQUOR FERRI PEPTONATI CUM MANGANO — Ferric Peptonate, 4.5, Soluble Manganese Citrate, 0.8, Ammonia Water (*U S P*), 1.3, Aromatic Elixir, 5.0, Alcohol (95 p c), 15.00, Distilled Water, *q s* to produce 100 — *U S N F*

Manganese Chloride, 0.35, Solution of Iron Peptonate, *q s* to produce 100 — *B P C*

FERRI ARSENAS.

IRON ARSENATE

ARSENATE OF IRON — *B P* '85

Fr., ARSFNIATE DE FER, GER., ARSENSAURES EISENOXYDUL,
SPAN., ARSENIATO DE HIERRO

A tasteless, olive-green, amorphous powder, consisting of Ferric Arsenate, $\text{Fe}_3(\text{AsO}_4)_2 \cdot 6\text{H}_2\text{O}$, eq. 550.12, Ferric Arsenate and some Iron Oxide, and ' ' ' ' not less than 12½ p c of hydrous, equivalent to 10 p c of anhydrous Ferrous Arsenate

Medicinal Properties — Similar to those of Arsenous Acid, the quantity of Iron in the dose is extremely small

Dose — $\frac{1}{8}$ to $\frac{1}{4}$ grain = 0.004 to 0.016 gramme

Prescribing Note — *Best given in pill well triturated with Milk Sugar, and massed with a little Glucose*

Antidotes — *See Acidum Arseniosum*

Not Official — *Mistura Ferri Arsenicalis, Pilula Ferri Arsenicalis, Ferri Aisenio Citras Ammoniata, Injectio Ferri Aisenatis Solubilis*

Foreign Pharmacopœias — Official in Fr, Mex (*Aiseniato de Fierro*) and Span Not in the others

Tests — Iron Arsenate dissolves in Hydrochloric Acid, and the solution answers the tests distinctive of both Ferrous and Ferric salts given under Ferrum After separation of the Iron, the neutralised filtrate should yield a reddish-brown precipitate with Silver Ammonio-nitrate Solution, and a white crystalline precipitate with Magnesium Ammonio-sulphate It is officially required to indicate 10 p c of anhydrous or nearly 12½ p c of hydrous Ferrous Arsenate as determined by titration with Volumetric Potassium Bichromate Solution, using Potassium Ferricyanide Solution as an indicator, 1 gramme requiring at least 6.7 c c The solution in Hydrochloric Acid should yield no decided turbidity with Barium Chloride Solution, indicating the absence of more than a trace of Sulphates

Not Official

MISTURA FERRI ARSENICALIS — Arsenical Solution, 2 minims, Iron and Ammonium Citrate, 5 grains, Tincture of Calumba, 10 minims, Water, to 1 fl oz — *St Thomas's*

This has been incorporated in the *B P C*

Citrate of Iron and Ammonium, 8 grains, Arsenical Solution, 5 minims, Tincture of Calumba, 30 minims, Water, to 1 fl oz — *University*

Arsenical Solution, 5 minims, Iron and Ammonium Citrate, 6 grains, Infusion of Quassia, to 1 fl oz — *Guy's*

PILULA FERRI ARSENICALIS — Arsenious Anhydride, $\frac{3}{16}$ grain, Excised Ferrous Sulphate, 3 grains, Excipient, *q s* for one pill — *University*
Arsenious Acid, $\frac{1}{10}$ grain, Excised Ferrous Sulphate, 3 grains, Milk Sugar and Syrup of Glucose, *q s* for one pill — *B P C*

FERRI ARSENIO-CITRAS AMMONIATA — Green or yellowish green deliquescent scales, containing 1.4 p c Arsenious Acid and 15 to 18 p c of Iron Readily soluble in Water A valuable antiperiodic Best administered by subcutaneous injection

INJECTIO FERRI ARSENATIS — A neutral, sterilised solution, containing 2.5 p c of the above salt, specially prepared for hypodermic administration The dose, which is 1 c c, contains 0.0035 gramme Arsenious Acid and from 0.00375 to 0.0045 gramme of Iron

FERRI CACODYLAS — *See under Sodii Cacodylas*

Not Official

FERRI BROMIDUM

The commercial salt is in greyish white crystalline masses, coated with red insoluble Oxybromide, which amounts to about 0.5 p c

It generally contains about 18 p c of Water, corresponding nearly with the formula $\text{FeBr}_2 \cdot 3\text{H}_2\text{O}$, eq 347.29 When this is not allowed for, a Syrup or Liquor made from the solid Bromide, and calculated as if anhydrous, will be proportionately weaker than when made from Iron Wire

Foreign Pharmacopœias —Not in any

LIQUOR FERRI BROMIDI FORTIS —A clear green liquid Sp gr 1.554

Each fl dr̄m contains 36 grains of Iron Bromide FeBr_2 214.9)

This solution keeps well in a corked bottle, with Wire immersed in it, and on filtration gives a clear green liquid

With the addition of a small quantity of Hypophosphorous Acid, the Liquor will keep very well

Foreign Pharmacopœias —Official in Mex (Bromuro Ferroso) and Port (Brometo Ferroso), both solid, no solution Not in the others

SYRUPUS FERRI BROMIDI —Strong Solution of Iron Bromide (filtered), 1, Simple Syrup, 7, mix

Contains $4\frac{1}{2}$ grains of Iron Bromide in each fl dr̄m

Medicinal Properties —A tonic in anæmia and amenorrhœa

Syrupus Ferri Bromidi —Iron Wire free from oxide, 2.5, Bromine, 6, Refined Sugar, 70, Distilled Water, q s to yield 100 —B P C

It is not stated in the B P C whether the Bromine is by weight or measure, but as it is taken from the *Conference* formula, it is most probably by weight, and rather less Bromine is used in the B P C It was subsequently stated to be by weight

Each fl dr̄m contains about $4\frac{1}{2}$ grains of Ferrous Bromide

Dose —30 to 60 minims = 1.8 to 3.6 cc

SYRUPUS FERRI BROMIDI CUM QUININA —Acid Quinine Hydrobromide, $\frac{1}{2}$ oz, Diluted Hydrobromic Acid, 2 fl dr̄m, Distilled Water, 1 fl oz, dissolve the Quinine salt in the Acid and Water mixed, then add Syrup of Ferrous Bromide, q s to yield 12 $\frac{1}{2}$ fl oz —B P C

Dose —30 to 60 minims = 1.8 to 3.6 cc

1 fl dr̄m contains about $1\frac{1}{10}$ grains of Acid Quinine Hydrobromide, and about 4 grains Ferrous Bromide

It is rather stronger in Quinine than the *Conference* formula

SYRUPUS FERRI BROMIDI CUM STRYCHNINA —Strychnine 1 grain, Diluted Hydrobromic Acid, 70 minims, Syrup Ferrous Bromide, to 8 fl oz

60 minims contains $\frac{1}{10}$ grain of Strychnine

SYRUPUS FERRI BROMIDI CUM QUININA ET STRYCHNINA —Dissolve 1 grain of Strychnine, in powder, in 8 fl oz of the Syrup of Ferrous Bromide with Quinine given above —B P C

1 fl dr̄m contains about $\frac{1}{10}$ grain Strychnine, about $1\frac{1}{10}$ grains Quinine Acid Hydrobromide and about 4 grains Ferrous Bromide

Dose —30 to 60 minims = 1.8 to 3.6 cc

FERRI CARBONAS SACCHARATUS.

SACCHARATED IRON CARBONATE

FR, SACCHARURE DE CARBONATE FERREUX, GER, ZUCKERHALTIGES FERRO CARBONAT

Dull, greyish-brown, amorphous, odourless at first a sweet and subsequently a ferruginous taste stated to consist of Ferrous Oxy carbonate, $x\text{FeCO}_3, y\text{Fe(OH)}_2$, in a greater or less degree of oxidation, mixed with Sugar, the Ferrous salt, if reckoned as Ferrous Carbonate, FeCO_3 , eq 115.15, 33 $\frac{1}{3}$ p c of the mixture

A new method of preparing Saccharated Carbonate of Iron is recommended by Mr J H Franklin, he proposes the use of liquid Glucose instead of Sugar, the percentage of ferrous carbonate obtained is about double that obtained by the official process, and keeps perfectly in a well-closed bottle. It is useful in the preparation of pills, tablets and capsules—*P J '07*, n 114, 155

Medicinal Properties—An excellent chalybeate, readily taken and well borne. Not astringent. Useful in anæmia, and in anæmic forms of amenorrhœa, neuralgia and sciatica. Ferrous Carbonate, in the form of 'Blaud's Pills,' is a very popular medicine.

Dose—10 to 30 grains = 0.65 to 2 grammes

The above dose is equivalent to $3\frac{1}{2}$ to 10 grains = 0.216 to 0.65 gramme of Ferrous Carbonate

Prescribing Notes—*Given in cachets, lozenges, or pills. Sometimes ordered in the form of Powders to be taken on bread and butter. A good pill can be made by adding Dispensing Syrup q s. It can also be taken as an effervescent granule.*

Incompatibles—Acids and Acidulous salts, all Vegetable astringents

Official Preparations—*Mistura Ferri Composita and Pilula Ferri.* Although not actually prepared from the Saccharated Iron Carbonate, they are here grouped for comparison.

Not Official—*Massa Ferri Carbonatis, Pilulæ Ferri Carbonatis, Trochisci Ferri Carbonatis Saccharati, Ferri Oxidum Saccharatum*

Foreign Pharmacopœias—Official in Austr and Swiss (*Ferrum Carbonicum Saccharatum*), contains about 20 p.c. of Carbonate, Belg (*Carbonas Ferri Saccharatus*), 20 p.c., U.S. contains 15 p.c., Ger, Jap and Russ, 9.5 to 10 p.c. of Iron equal to about 20 p.c. of Carbonate, Norw (*Hydratocarbonas Ferrosus Saccharatus*). No Sugar Jap (*Ferrum Subcarbonicum*), and Mex (*Carbonato de Hierro*). Not in the others.

Tests—Saccharated Iron Carbonate dissolves with effervescence in diluted Hydrochloric Acid, and the solution yields with Potassium Ferrocyanide or Potassium Ferricyanide Solution a blue precipitate. It is officially required to contain about $33\frac{1}{2}$ p.c. of the Ferrous Salt if reckoned as Ferrous Carbonate, the *U.S.P.* preparation is required to contain not less than 15 p.c. of Ferrous Carbonate, and the *P.G.* from 9.5 to 10 p.c. of Iron, corresponding to 19.7 to 20.7 p.c. of Ferrous Carbonate. The *B.P.* employs Volumetric Potassium Bichromate Solution for the determination, and Potassium Ferricyanide Solution as an indicator, dissolving the Carbonate in warm concentrated Phosphoric Acid, notwithstanding it having been shown that warming on a water-bath even for 10 minutes introduced an error of 25 p.c. The *U.S.P.* dissolves the Carbonate in Diluted Sulphuric Acid and performs the titration with Tenth-normal Volumetric Potassium Dichromate Solution. The *P.G.* converts the whole of Ferrous salt into the Ferric condition, and determines the total Ferric Iron with Potassium Iodide Solution, titrating the liberated Iodine with Tenth-normal Volumetric Sodium Thiosulphate Solution.

A 2 p.c. solution of the Carbonate in sufficient Hydrochloric Acid to effect solution and ensure a slight excess of acid should yield no pronounced turbidity with Barium Chloride Test Solution.

Barium Nitrate (or Chloride)—A solution of Saccharated Iron Carbonate prepared as above should not give more than a slight cloudiness with T S of Barium Chloride, *U S P*. The *P G* requires that a solution of the Saccharated Iron Carbonate in Water (1-50) obtained by means of the least possible quantity of Hydrochloric should only be rendered slightly turbid by T S of Barium Nitrate, *P G*.

Volumetric Determination—The solution of 1 gramme of Saccharated Iron Carbonate in excess of warm concentrated Phosphoric Acid diluted with Water should require at least 29 c c of the Volumetric of Potassium Dichromate Solution, using Potassium Ferricyanide Solution as an indicator, *B P*, the solution obtained by dissolving 1.15 gramme in 10 c c of dilute Sulphuric Acid, diluted to about 100 c c with Water should require not less than 15 c c of Tenth-normal Volumetric Solution of Potassium Dichromate for complete titration, using Potassium Ferricyanide Solution as an indicator, *U S P*, the *P G* requires that 1 gramme be dissolved in 10 c c of dilute Sulphuric Acid without heat. To this is added solution of Potassium Permanganate (5-1000) until the faint transitory reddening just becomes permanent, then 2 grammes Potassium Iodide. The mixture is allowed to stand for one hour in a closed vessel at ordinary temperatures, and then titrated with Tenth-normal Volumetric Solution of Sodium Thiosulphate, for combination with the free Iodine 17 to 17.8 c c of Tenth-normal Volumetric Solution of Sodium Thiosulphate should be necessary.

Preparations

MISTURA FERRI COMPOSITA. COMPOUND MIXTURE OF IRON. *N O Syn*—GRIFFITH'S MIXTURE

Reduce 60 grains of Myrrh to powder, and mix it with 30 grains of Potassium Carbonate and 60 grains of Refined Sugar, form this into a smooth thin paste, by rubbing with a small quantity of Rose Water. Gradually add more Rose Water and 50 minims of Spirit of Nutmeg until the product measures 7 fl oz. Dissolve 25 grains of Ferrous Sulphate in 3 fl oz of Rose Water, and mix with the above.

It is convenient to keep the first part of the mixture ready made, and to add the Ferrous Sulphate solution when required for use.

Dose— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 c c

The following modification has been suggested by Mr J H Franklin—Saccharated Carbonate of Iron (with Glucose), 16 grains, Syrup of Glucose, 3 fl drms, Gum Acacia (in powder), 20 grains, Tincture of Myrrh, 4 fl drms, Spirit of Nutmeg, 50 minims, Rose Water to produce 10 fl oz. Reduce the Saccharated Carbonate of Iron to a fine powder, triturate with the Syrup of Glucose and continue the trituration with a few drops of Rose Water to form a smooth thin paste, gradually add more of the Rose Water, and add the Acacia diffused in the Tincture of Myrrh and Spirit of Nutmeg, finally making the product measure 10 fl oz with Rose Water.—*P J '07*, n 155, *C D '07*, n 180.

This has been incorporated in the *B P C* under the title **Mistura Ferri Carbonatis Composita**.

Foreign Pharmacopœias—Official in Dan, similar to Brit, but with three times as much Sugar, and without Nutmeg, Norw, without Nutmeg, with Peppermint Water, Swed, Emulsio Myrrhæ Ferratæ, with Peppermint Water and Tincture of Lavender in the place of Rose Water and Nutmeg, *U S* similar to Brit, but with Spirit of Lavender in the place of Nutmeg. Not in the others.

PILULA FERRI. IRON PILL

Mix 150 grains of Syrup, 10 grains of Glycerin and 20 grains of Distilled Water, with this incorporate 150 grains of Exsiccated Ferrous Sulphate, add 95 grains of Exsiccated Sodium Carbonate,

and mix quickly Allow 15 minutes for the salts to react, and make into a pill mass by the addition of 50 grains of powdered Gum Acacia and 15 grains of powdered Tragacanth

If divided into 5 grain pills, each pill will contain about 1 grain of Ferrous Carbonate

Dose—5 to 15 grains = 0.32 to 1 gramme

Pilula Ferri Carbonatis (*B P '85*)—Made with Saccharated Iron Carbonate, 4, Confection of Roses, 1, contains rather more Ferrous Carbonate than *Pilula Ferri* (*B P '98*)

The following modification has been suggested by Mr J H Franklin—Saccharated Carbonate of Iron (with Glucose), 648 grains, Liquorice Root (in powder), 162 grains, Liquid Glucose, 216 grains, Water, 54 grains Make a mass, and divide into pills weighing $2\frac{1}{2}$, 5 and $7\frac{1}{2}$ grains each—*P J '07*, 11 115

This has been incorporated in the *B P C*

Vallet's mass is made by precipitating and washing the Iron Carbonate, and mixing it with Honey and Milk Sugar to form a mass See below

Blaud's Pills are made by mixing (in the pill mass) dried Ferrous Sulphate and dried Potassium or Sodium Carbonate See below

Foreign Pharmacopœias—Official in Belg and Dutch (*Pilulæ Bland*), Dan and Norw (*Pilulæ Blandii*) also (*Pilulæ Ferri Compositæ*), Fr (*Pilules de Carbonate Ferreux*, formule de Vallet, and *Pilules de Carbonate de Fer composées*, *Pilule de Bland*, Ger and Jap (*Pilulæ Ferri Carbonici Blandii*), Austri (*Pilulæ Ferri Carbonici*), Ital (*Pillule di Carbonato Ferroso*) (*Pillule di Bland*) also (*Pillule di Vallet*) Mex (*Pildoras de Bland* and *Pildoras de Vallet*), Port (*Pilulas de Carbonato Ferroso*), Span (*Pildoras de Bland*), Swed (*Pilulæ Ferratæ Blandii* and *Pilulæ Myrihæ Ferratæ*), Swiss (*Pilulæ Ferratæ Blandii* and *Pilulæ Ferri Carbonici*) (*Pil Valletii*), US (*Pilulæ Ferri Carbonatis*) (*Blaud's Pills*), also (*Massa Ferri Carbonatis*) (*Vallet's Mass*) Not in the others

Not Official

MASSA FERRI CARBONATIS (*Vallet's Mass*)—Dissolve 100 of Ferrous Sulphate and 46 of Monohydrated Sodium Carbonate, each separately, in 200 of boiling Distilled Water, and, having added 20 of Syrup to the solution of the Iron salt, filter both solutions and allow them to become cold, gradually add the Iron solution to the Sodium solution in a 500 bottle, rotating it until Carbonic Acid gas no longer escapes Add Distilled Water, *q s* to fill the bottle, then cork it and set aside so that the Ferrous Carbonate may subside Pour off the supernatant liquid, and wash the precipitate with a mixture of Syrup 1, Water 19, by decantation until the washings no longer have a saline taste Drain and press, mix the precipitate at once with 38 of Clarified Honey and 25 of Sugar, and evaporate the mixture in a tared dish on a water-bath, with constant stirring, until it is reduced to 100—*U S*

PILULÆ FERRI CARBONATIS (*Blaud's Pills*)—Rub 8 grammes of Potassium Carbonate in a mortar with about 10 drops each of Glycerin and Water, then add 16 grammes of Ferrous Sulphate and 4 grammes of Sugar, previously triturated together to a uniform powder, and rub the mass thoroughly until it assumes a greenish colour When the reaction has terminated incorporate 1 gramme of Tragacanth and 1 gramme of Althæa, and, if necessary, a little more Water, so as to obtain a mass of pillular consistence Divide this into 100 pills—*U S*

TROCHISCI FERRI CARBONATIS SACCHARATI—Containing 3 grains of Saccharated Carbonate in each

Dose—1 to 3 lozenges

FERRUM OXIDUM SACCHARATUM (*Ger and Austri*)—A reddish-brown powder, with a sweet, slightly ferruginous taste, a mixture of Hydrated Ferric Oxide and Sugar, containing the equivalent of 2.8 p.c of Iron

Dose—5 to 15 grains = 0.32 to 1 gramme

FERRI ET AMMONII CITRAS.

IRON AND AMMONIUM CITRATE

Thin, translucent, deep ruby-red, odourless, deliquescent scales, with a ferruginous and somewhat astringent taste

Solubility—10 in 5 of Water, and measures $10\frac{1}{2}$, 2 dissolved in 3 of Water measure 4, almost insoluble in Alcohol (90 p c)

Medicinal Properties.—As a hæmatinic, it is a very effectual salt, and it possesses scarcely any astringency or tendency to cause gastric irritation or constipation, it may often be given when the stomach will not bear the more astringent preparations of Iron. It becomes moist if kept in paper.

A useful prescription for combating the anæmia which is often a marked feature of neurasthenia is Ferric Ammonium Citrate, 7 grains, Liquor Arsenicalis, 5 minims, Potassium Bromide, 10 grains, Liquor Ammon Acet, 1 drim, Chloroform Water, to 1 oz.—*B M J '06*, 1 494

Dose.—5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Note—*Generally prescribed in solution with Tincture of Orange, which covers the taste well*

An Aqueous Solution, 2 fl oz representing 480 grains of the scale preparation, is convenient for dispensing, and keeps well

Incompatibles—Mineral Acids, Vegetable astringents, and fixed Alkalis

Official Preparation—Vinum Ferri Citratis

Not Official—Mistura Ferri cum Ammonia

Foreign Pharmacopœias—Official in U.S., Jap and Swiss (Ferrum Citricum Ammoniatum), Belg (Ferrum Citricum), Fr (Citrato de Fer Ammoniacal), Ital (Citrato di Ferro Ammoniacale), Mex (Citrato de Fierro Amoniacal), Norw (Citrato Ferrico-Ammonicus), Port (Citrato de Ferro Ammoniacal), Swed. (Citrato Ferricus), Swiss (Ferrum Citricum Ammoniatum), Span (Citrato Ferrico-Amónico) Not in the others. Ger has Ferrum Citricum Oxidatum

Tests.—Iron and Ammonium Citrate has a faintly acid reaction towards blue Litmus paper. When incinerated with free access of air it leaves a residue of 31 or 32 p c of Ferric Oxide. An aqueous solution when heated with an excess of Potassium Hydroxide Solution evolves the distinctive odour of Ammonia, and yields a brownish-red precipitate, and, if this precipitate be filtered off, the filtrate, when neutralised with Acetic Acid, yields on boiling with a little Calcium Chloride Solution a white precipitate.

Over and above the statement that when incinerated with free access of air it leaves 31 or 32 p c of Ferric Oxide, the *BP* gives no method for the detection of the Iron. The *USP* employs the Potassium Iodide process mentioned below, titrating the liberated Iodine with Tenth-normal Volumetric Sodium Thiosulphate Solution, using Starch Solution as an indicator. As thus determined, the percentage of metallic Iron should amount to not less than 16 p c, equivalent to not less than 22.8 p c of Ferric Oxide.

The more generally occurring impurities are fixed alkalis, Tartrates and Sulphates. Fixed Alkalis may be detected by the reaction

of the ash towards Litmus paper, it should not show an alkaline reaction towards red Litmus paper Tartrates are indicated by the appearance of a crystalline precipitate on the addition of an excess of Acetic Acid to the filtrate after removal of the Iron by boiling with Potassium Hydroxide Solution An aqueous solution should yield no pronounced turbidity with Barium Chloride Solution

Volumetric Determination—The *U S P* directs that 0.555 gramme of the salt be dissolved in 15 c.c. of Water and 2 c.c. of Hydrochloric Acid in a glass stoppered flask having a capacity of about 100 c.c., and 1 gramme of Potassium Iodide added The flask is then securely closed and the mixture kept at a temperature of 40° C (104° F) for half an hour and then cooled When titrated with Tenth-normal Volumetric Solution of Sodium Thiosulphate, using Starch T S as indicator, the mixture should require not less than 16 c.c. of the Volumetric Solution to discharge the colour of the liquid

Preparation

VINUM FERRI CITRATIS —WINE OF IRON CITRATE

Iron and Ammonium Citrate, 160 grains, Orange Wine, *q s* to yield 20 fl oz (1 grain in each fl dm)

Dose—1 to 4 fl drm = 3.6 to 14.2 c.c.

Official in Jap (*Vinum Ferri*), 1 in 50, **Mex** (*Vino de Fieiro*), 1 in 150, **Span** (*Vino Calibeado*), 1 and 200 of Malaga

Vinum Ferri (U S)—Iron and Ammonium Citrate, 4, Tincture of Sweet Orange Peel, 6, Syrup, 10, White Wine, *q s* to produce 100

Vinum Ferri Amarum, see p 517

Not Official

MISTURA FERRI CUM AMMONIA—Iron and Ammonium Citrate, 10 grains, Aromatic Spirit of Ammonia, 30 minims, Infusion of Quassia, to 1 fl oz—*King's*

Iron and Ammonium Citrate, 5 grains, Aromatic Spirit of Ammonia, 10 minims, Spirit of Chloroform, 5 minims, Infusion of Quassia, to 1 fl oz—*Royal Free*

FERRI ET QUININÆ CITRAS

IRON AND QUININE CITRATE

Thin, transparent, pale yellowish green, deliquescent scales possessing a bitter and ferruginous taste

It should be kept in well-closed vessels and protected as far as possible from the light

Solubility—2 in 1 of Water

Medicinal Properties—Bitter stomachic and tonic, combining the properties of both Iron and Quinine

63 grains contain 1 grain of Quinine

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes—Generally given in Mixture with Tincture of Orange and Spirit of Chloroform, or Syrup of Orange, or in Pills made with Alcohol (90 p.c.) *q s* It is sometimes prescribed with Potassium Citrate or Lithium Citrate, both of which have a tendency to throw out Quinine Citrate It can be given in the form of Effervescent Granules, dose one teaspoonful

For dispensing purposes, it is convenient to keep an aqueous solution, 2 fl oz = 480 g. ains of the salt

Incompatibles—Alkalis, then Carbonates and Citrates, Lithium Citrate, Tannic Acid, and vegetable astringents.

Not Official—Vinum Ferri Amatum, Vinum Ferri et Quininae, Ferri Quininae et Strychninae Citras, Ferri et Strychninae Citras

Foreign Pharmacopœias—Official in Austr (Ferrum Citricum Chinatum), Ger, Jap and Russ (Chininum Ferro-Citricum), Dan (Citras Ferricus cum Chinina), Noiw (Citras Ferricus cum Chinino) Port (Citrate de Ferro e de Quinina), Swed (Citras Ferrico Chinicus), Swiss (Chinino-Ferium Citricum), U S Not in the others U S has also Ferri et Quininae Citras Solubilis

Tests—Iron and Quinine Citrate dissolves readily in Water, yielding a solution which is very faintly acid in reaction towards blue Litmus paper. The aqueous solution yields with Potassium Hydroxide Solution a reddish-brown precipitate, and when heated evolves Ammonia (which fact is not noticed in *BP*), with Ammonia Solution it yields a white curdy precipitate, with Potassium Ferrocyanide and with Potassium Ferricyanide blue precipitates, with Tannic Acid a bluish black precipitate.

It is officially required to contain 15 p c of Ether-soluble alkaloid, which when neutralised by Sulphuric Acid should answer to the tests for Quinine Sulphate. No standard for the percentage of Iron is given. The *USP* requires the salt to contain not less than 11.5 p c of dried Quinine, and Ferric Citrate corresponding in amount to not less than 13.5 p c of metallic Iron. The *PG* preparation contains from 9 to 10 p c of Quinine, and is required to leave not less than 30 p c of Iron Oxide on ignition. An outline of the method adopted by the *BP* for the determination of the Quinine is given below. The extracted alkaloid is required to be almost entirely soluble in a little purified Ether, to leave but a minute residue on ignition, and, when neutralised by Sulphuric Acid, to answer the tests distinctive of Quinine Sulphate. The *USP* employs Chloroform as a solvent for the Quinine, and requires that the dried residue should conform to the reactions and tests for Quinine. The Iodometric method is adopted for the determination of the Iron, the determination being conducted on the liquid remaining after the removal of the Quinine. The *PG* employs Ether as a solvent in the Quinine determination. Allon has pointed out that in shaking out with Chloroform or Ether a considerable excess of Ammonia should be present, and the volume of the solvent should equal that of the ammoniacal liquid. The alkaloidal residue should be dried at 110° to 120° C (230° to 248° F) a constant weight being difficult to obtain at a water-bath temperature.

The more generally occurring impurities are fixed alkalis, which may be detected by the alkaline reaction of the residue left on ignition, and Tartrates, which yield a crystalline precipitate, when Acetic Acid is added in slight excess to the filtrate after removal of the Iron by precipitation with boiling Potassium Hydroxide Solution.

Gravimetric Determination—Dissolve a weighed quantity of 5 grammes of the salt in 45 c c of Water, add Ammonia Solution in slight excess, extract

the liberated alkaloid by repeated shakings with Ether. Separate the ethereal solutions, mix, evaporate to dryness, and when completely dried at 120° C (248° F), cool and weigh. The residue should amount to 0.75 gramme, *B P*, a weighed quantity of 1.11 gramme of the salt is dissolved in 20 c c of Water, transferred to a separator, rendered alkaline with 5 c c of Ammonia Solution, and the mixture shaken out for 1 minute with 10 c c of Chloroform. The chloroformic layer is separated and the agitation twice repeated with successive quantities each of 10 c c of Chloroform. The Chloroform solutions are mixed, transferred to a tared dish, the Chloroform evaporated spontaneously, the residue dried at 100° C (212° F) till constant in weight. It should weigh not less than 0.1276 gramme, which is equivalent to at least 11.5 p c of Quinine. The aqueous liquid from the above determination is freed from Chloroform by heating on a water bath until all ammoniacal and chloroformic odours have disappeared, cooled, and diluted with Water to 50 c c. A measured quantity of 25 c c is transferred to a glass stoppered flask capable of holding about 100 c c, 3 c c of Hydrochloric Acid and 1 gramme of Potassium Iodide added, the flask securely stoppered, and the mixture allowed to stand half an hour at 40° C (104° F). When cool not more than 13.5 c c of Tenth normal Volumetric Sodium Thiosulphate shall be required to discharge the colour of the liquid, Starch Solution being employed as an indicator. 1 c c of Tenth normal Volumetric Sodium Thiosulphate Solution indicating 1 p c of metallic Iron, *U S P*, a weighed quantity of 1 gramme of the salt is dissolved in 4 c c of Water, and sufficient Sodium Hydroxide Solution (15 p c) added to ensure a strongly alkaline reaction. The mixture is then shaken out three times in succession with 7 c c of Ether. The separated ethereal layers are mixed, evaporated to dryness, and the residue dried at 100° C (212° F). It should weigh at least 0.09 gramme, *P G*.

Not Official

VINUM FERRI AMARUM—Soluble Iron and Quinine Citrate, 5, Tincture of Sweet Orange Peel, 6, Syrup, 30, White Wine, *q s* to produce 100—*U S P*

VINUM FERRI ET QUININÆ—Iron and Quinine Citrate, 2, Detanated Sherry, *q s* to produce 100—*B P C*

Ferri, Quininæ et Strychninæ Citras, resembling the above but containing in addition 1 p c of Strychnine, and **Ferri et Strychninæ Citras** (*U S*), similar to the above but without Quinine, are both scale preparations, the doses of which are 2 to 5 grains = 0.13 to 0.82 gramme

Not Official

FERRI HYPOPHOSPHIS

There are two Iron Hypophosphites, the Ferrous and the Ferric. The latter is used in most of the American and other proprietary Syrups of the Hypophosphites. The Ferric salt has now replaced the Ferrous salt in the *B P C* preparations.

FERROUS HYPOPHOSPHITE, when freshly prepared, is a greenish crystalline powder, soluble about 1 in 10 of Water, but the commercial salts are so insoluble as to be practically useless for pharmaceutical purposes.

FERRIC HYPOPHOSPHITE—This compound is obtained as a white precipitate on adding a solution of a soluble Hypophosphite to one of Ferric Chloride containing as little free acid as possible.

It is fairly insoluble in Water, but with the addition of Potassium Citrate it dissolves readily to a green solution, which forms with Sugar a pale yellow *neutral* Syrup, permanent and unalterable by exposure to air, which may be combined with other soluble Hypophosphites, Quinine Hydrochloride, and Strychnine without the addition of acid, and is free from all the pharmaceutical objections attaching to Hypophosphite Syrups containing Iron in the ferrous condition.

Official in U.S.

It is usually sold as **Compound Syrup of Hypophosphites**, and is also made without Quinine to which it is added, who are peculiarly susceptible to that drug, it is then prescribed 'sine Quinina'.

LIQUOR FERRI HYPOPHOSPHITIS FORTIS — Solution of Ferric Sulphate, 14 2, Solution of Ammonia, 28, Citric Acid, 7 6, Sodium Hypophosphite, 9 6, Sodium Citrate, 6 6, Distilled Water *q s* and Chloroform Water (1 in 200) *q s* to produce 100 — *B P C*

This formula was devised (*Y B P* '07, 265) as an improvement on the old *B P C* method

The solution of Ferric Sulphate is diluted with an equal volume of Water and added to the solution of Ammonia also diluted with an equal volume of Water. After the precipitated Ferric Hydroxide has subsided sufficiently, wash it by decantation with Distilled Water until free from Sulphates, collect the precipitate and drain it, dissolve it in the Citric Acid previously dissolved in 20 c of Distilled Water by the aid of a water-bath. When the solution is clear add the Sodium Hypophosphite and continue the heat until a clear greenish solution is produced, then add Sodium Citrate, filter and pass sufficient Chloroform Water through the filter to make up 100 of product.

The *B P C Supplement* replaces the 14 2 of solution of Ferric Sulphate by 12 9 of Ferric Citrate.

LIQUOR HYPOPHOSPHITUM — Calcium Hypophosphite, 3 5, Sodium Hypophosphite, 2, Potassium Hypophosphite, 1 75, Citric Acid, 1 6, Water, *q s* to produce 100, dissolve and filter — *U S N F* 1896

This has been incorporated in the *B P C*

In *U S N F* 1906 the 1 6 of Citric Acid is replaced by 0 6 of Hypophosphorous Acid, *U S P*

Dose — 10 to 30 minims = 0 6 to 1 8 c c

LIQUOR HYPOPHOSPHITUM COMPOSITUS *Syn.* Liquor Hypophosphitis Compositus — Dissolve 320 grains of Calcium Hypophosphite, 320 grains of Sodium Hypophosphite, and 160 grains of Magnesium Hypophosphite in 12 fl oz of Distilled Water, add 6 fl oz of Strong Solution of Ferric Hypophosphite, filter, and add Distilled Water to make the product 20 fl oz — *B P C Formulary* 1901

Each fl drim = 2 grains each of Sodium and Calcium Hypophosphites, 1 grain Magnesium Hypophosphite, and 1½ grains of Ferric Hypophosphite

Dose — ½ to 2 fl drim = 1 8 to 7 1 c c

This has been incorporated in the *B P C* as follows — Calcium Hypophosphite, 8 5, Magnesium Hypophosphite, 1 75, Sodium Hypophosphite, 3 5, Strong Solution of Ferric Hypophosphite, 30, Distilled Water, sufficient to produce 100

SYRUPUS FERRI HYPOPHOSPHITIS — Strong Solution of Ferric Hypophosphite, 1, Syrup, 4 — *B P C Formulary* 1901, incorporated in the *B P C*

Each fl drim = about 1 grain of Ferric Hypophosphite

Dose — ½ to 2 fl drim = 1 8 to 7 1 c c

SYRUPUS HYPOPHOSPHITUM — Calcium Hypophosphite 4 5, Potassium Hypophosphite, 1 5, Sodium Hypophosphite 1 5, Distilled Hypophosphorous Acid, 0 20, Tincture of Fresh Lemon Peel, 0 5, Sugar, 65, Water, *q s* to make 100 — *U S P*

Dissolve the Hypophosphites in 45 of the Water, add the Tincture and the Acid, filter, and in the filtrate dissolve the Sugar without heat, make up to 100 with Water

This has been incorporated in the *B P C*

SYRUPUS HYPOPHOSPHITUM COMPOSITUS — Calcium Hypophosphite, 80 grains, Magnesium Hypophosphite, 40 grains, Potassium Hypophosphite, 40 grains, Citric Acid, 20 grains, Distilled Water, 8 fl oz of Chloroform Water, and 1 gr of Strychnine dissolved in 1 fl drim of Hypophosphorous Acid, and then 1 fl oz of Strong Solution of Ferric Hypophosphite, add 14 oz of Refined Sugar and dissolve without heat, add sufficient Chloroform Water to make 20 fl. oz. and strain through flannel. — *B. P. C.*

Each fl dr̄m contains $\frac{1}{150}$ grain Strychnine, and $\frac{1}{2}$ grain of Quinine Hypo phosphite

Dose — $\frac{1}{2}$ to 2 fl dr̄m = 18 to 71 cc

The employment of Potassium Citrate (1.033 gramme) in the place of Ammonium Citrate for dissolving the precipitated Ferric Hypophosphite, as originally recommended in the *Companion*, is also advocated — *P J '02*, 11 532

Syrupus Hypophosphitum Compositus — Rub 2.25 of Ferric Hypo phosphite and 2.25 of Manganese Hypophosphite with 3.75 of Sodium Citrate, add 30 cc of Water and warm the mixture for a few minutes, until a clear greenish solution is obtained. Dissolve 35 of Calcium Hypophosphite, 17.5 of Potassium Hypophosphite in a mixture of 450 cc of Water, and 5 cc of Diluted Hypophosphorous Acid, then dissolve 1.10 of Quinine and 0.115 of Strychnine in a mixture of 30 cc of Water and 10 cc of Diluted Hypophosphorous Acid, mix the solutions and finally dissolve in them 775 of sugar. Strain the Syrup, if necessary, and add Water *q s*, through the strainer, to produce 1000 cc — *U S*

Not Official FERRI IODIDUM

IRON IODIDE

FeI_2 , eq 307.40

In reddish brown, deliquescent dense masses, easily soluble in Water, leaving only a slight residue, and forming a reddish yellow solution owing to partial oxidation. The solution may be made green by either hot or cold digestion over bright Iron Wire.

Medicinal Properties — It combines the properties both of Iodine and Iron, and is a most valuable tonic and alterative in the treatment of scrofulous and syphilitic diseases.

Prescribing Notes — *Best given in the form of the official Syrup of Ferrous Iodide, it is also given in the form of pills massed with powdered Gum Acacia and Dispensing Syrup, q s. In some cases Liquorice Powder must be used instead of Dispensing Syrup.*

Official Preparation — Syrupus Ferri Iodidi

Foreign Pharmacopœias — Official in Mex (Yoduro Ferroso) and Port. Not in the others. Jap has Ferrum Iodati Saccharatum.

LIQUOR FERRI IODIDI FORTIS — A clear, greenish liquid.

Each fl dr̄m contains 44 grains of Ferrous Iodide ($\text{FeI}_2 = 307.40$). It can be diluted 1 to 7 with Syrup to prepare a Syrup of Iron Iodide, or with Water to make Liquor Ferri Iodidi the same strength as the Syrup.

With the addition of a small quantity of Hypophosphorous Acid, the solution will keep well for a long time, but in this case, when diluting with Syrup, it must be remembered that the official Syrup does not contain Hypophosphorous Acid.

The *B P C* gives a formula for the strong Liquor containing 7.5 pc of Diluted Hypophosphorous Acid (*U S P*).

Foreign Pharmacopœias — Official in Belg, Ger (Liquor Ferri Iodati), Russ and Swiss (Ferrum Iodatum Solutum), all contain 50 pc of Ferrous Iodide, Mex (Solucion oficial de Yoduro ferroso) 20 pc.

Incompatibles — Acids, Acid salts, Alkalis and their Carbonates, Lime Water, vegetable astringents.

PILULÆ FERRI IODIDI (U S) — Reduced Iron, 4 grammes, Iodine, 5 grammes, Glycyrrhiza, in No. 60 powder, 4 grammes, Sugar, in fine powder, 4 grammes, Extract of Glycyrrhiza, in fine powder, 1 gramme, Acacia, in fine powder, 1 gramme, Water, a sufficient quantity. To the Reduced Iron, contained in a small mortar, add 6 cc of Water, and then gradually the Iodine, constantly

tritulating, until the mixture ceases to have a reddish tint. Then add the remaining powders previously well mixed together, and mix the whole thoroughly. Transfer the mass to a porcelain capsule, and evaporate the excess of moisture, on a water-bath, with constant stirring, until the mass has acquired a pilular consistence. Coat with Balsam of Tolu dissolved in Ether. To make 100 pills.

Pilula Ferri Iodidi was official in *B P* '85, but omitted in 1898. It has been incorporated in the *B P C*.

Foreign Pharmacopœias—Official in Belg., Dan., Dutch, Fr., Ital., Mex., Norw., Port., Span., Swed. and Swiss, each pill contains about $\frac{3}{4}$ grain Iodide of Iron. Hung., about 1 grain, and all coated with Balsam of Tolu dissolved in Ether, except Dutch, which uses Tolu in Chloroform, and Swiss, not coated. Not in the others.

Official Preparation

SYRUPUS FERRI IODIDI. SYRUP OF FERROUS IODIDE

Iron Wire, $\frac{1}{2}$ oz., Iodine, 726 grains, Refined Sugar, $16\frac{1}{2}$ oz., Distilled Water, *qs* to produce 20 fl. oz. of a pale green Syrup containing 3 grains of Ferrous Iodide in 33 minims.

Dose—30 to 60 minims = 1 S to 3 C c.

This Syrup is very liable to become discoloured. It may be due to one or other of two causes: (1) Oxidation of Iron, which may be prevented by careful manipulation, or removed by Hypophosphorous Acid. (2) Slight caramelisation of the sugar by overheating, this cannot be removed by reducing agents.

Foreign Pharmacopœias—Official in Brit., 9.83 to 10 p.c. of Iron Iodide, Austr., Belg., Dan., Dutch, Ger., Jap., Russ., Span., Swiss and U.S., 5 p.c., Fr. and Port., 0.5 p.c., Hung., 12.2 p.c., Ital., 0.6 p.c., Mex., 1 p.c., Norw. and Swed., 10 p.c. All by weight.

The Brussels Conference agreed to a strength of 5 p.c. anhydrous Ferrous Iodide.

Tests.—Syrup of Ferrous Iodide has a specific gravity of 1.380 to 1.387, the *U.S.P.* syrup a specific gravity of about 1.349 at 25° C (77° F). When diluted with Water it affords with Potassium Ferricyanide Solution a dark blue precipitate. The diluted syrup mixed with Starch Solution yields on the addition of a little Chlorine Water a deep blue coloration. It is officially required to contain not less than 9.83 p.c. w/v, equivalent to 7.1 p.c. w/w, nor more than 10.14 p.c. w/v, equivalent to 7.31 p.c. w/w, of anhydrous Ferrous Iodide, as determined by decomposition of the Ferrous Iodide with Sodium Carbonate and titration of the exactly neutralised solution of Sodium Iodide with Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator. The *U.S.P.* syrup is required to contain about 5 p.c. w/w, equivalent to 6.74 p.c. w/v of Ferrous Iodide. The Ferrous Iodide is estimated by indirect titration with Tenth-normal Volumetric Silver Nitrate Solution, the excess of Silver being titrated with Tenth-normal Volumetric Potassium Sulphocyanate Solution, Ferric Ammonium Sulphate Solution being used as an indicator. The standard of 5 p.c. w/w of anhydrous Ferrous Iodide is that suggested by the Brussels Conference for the unification of the pharmacopœial formulas for potent drugs. Several processes claiming to be improvements on the official method of determination have been suggested but do not appear to have been adopted.

The *BPC* in an explanatory footnote to the preparation stated that it contained 1 p c of Ferrous Iodide. This was obviously incorrect, and has since been rectified.

Volumetric Determination—A measured quantity of 10 c c (equivalent to 18.87 grammes) of the Syrup is introduced into a flask of 100 c c capacity containing 1 gramme of dried Sodium Carbonate dissolved in 10 c c of Water. The mixture is shaken until complete interaction has taken place, sufficient Water is added to bring the volume of the liquid to 100 c c, the whole mixed and filtered. A measured quantity of 25 c c of the filtrate is exactly neutralised with diluted Nitric Acid and titrated with Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator. Not less than 16.0 nor more than 16.5 c c should be required. *BP* a weighed quantity of 10 grammes should be diluted with Water to 100 c c and 15.4 c c of the solution be mixed with 15 c c of Water, 6 c c of Tenth normal Volumetric Silver Nitrate Solution and 2 c c each of diluted Nitric Acid and Ferric Ammonium Sulphate Solution are added and the mixture thoroughly shaken. On titration with Tenth normal Potassium Sulphocyanate Solution not more than 1 c c should be required to produce a permanent reddish brown tint, 1 c c of Tenth normal Volumetric Silver Nitrate Solution corresponds to 1 p c of Ferrous Iodide, *USP*.

FERRI LACTAS

See under ACIDUM LACTICUM

Not Official

FERRI PERCHLORIDUM

FERRIC CHLORIDE

NO Syn—CHLORETUM FERRICUM

The commercial solid, or crystalline, Ferric Perchloride approximates to the formula $\text{Fe}_2\text{Cl}_6 \cdot 12\text{H}_2\text{O}$, eq. 586.90, it occurs in yellow, or yellowish-brown, crystalline masses, deliquescent in air. It is soluble in Water, Alcohol (90 p c), Ether and Glycerin.

Medicinal Properties—A powerful local styptic. 3 grains to an oz of Water for a **spray**, 60 to 120 grains to an oz of Water or Diluted Glycerin (Glycerin 1 and Water 1) for a **paint**.

2½ oz dissolved in 1 oz of Water makes a solution about the same strength as Liquor Ferri Perchlor. Fort.

For **Incompatibles**, see Tinctura Ferri Perchloridi.

The Anhydrous Ferric Chloride (Fe_2Cl_6 , eq. 322.84), prepared by sublimation, is in black metallic looking plates. It deliquesces rapidly on exposure to the air, and then solidifies again to a Hydrate ($\text{Fe}_2\text{Cl}_6 \cdot 12\text{H}_2\text{O}$), containing almost 40 p c of Water. Another Hydrate ($\text{Fe}_2\text{Cl}_6 \cdot 5\text{H}_2\text{O}$), containing 21.7 p c of Water (official in the Portuguese Pharmacopœia), can be obtained by evaporating an acid solution until syrupy, and then cooling it.

Foreign Pharmacopœias—Official in Austri. and Hung. (Ferrum Sesquichloratum Crystallisatum), Dan., Dutch, Norw. and Swed. (Chloretum Ferricum), Mex. (Cloruro Ferrico), Port. (Chloreto Ferrico Anhydrido), also Crystallizado, Ger., Jap. and Russ. (Ferrum Sesquichloratum), Span. (Cloruro Ferrico) (Anhydrous and the Hydrate), Swiss (Ferrum Sesquichloratum), U.S. (Ferri Chloridum).

FERRI PERCHLORIDI LIQUOR FORTIS.

STRONG SOLUTION OF FERRIC CHLORIDE

A dark reddish-brown liquid, possessing a powerfully styptic taste, readily miscible with Water, and Alcohol (90 p c)

Medicinal Properties—A powerful local styptic and astringent, escharotic. The more dilute forms are used internally to arrest hæmorrhage in the gastro-intestinal or urinary tracts. See also 'Tinctura Ferri Perchloridi,' p 524

The liquor (not fortis) in $\frac{1}{2}$ drm doses thrice daily for hæmorrhagic gastric oozing—*L* '06, ii 1189

Official Preparations—Liquor Ferri Perchloridi and Tinctura Ferri Perchloridi

Not Official—Glycerinum Ferri Perchloridi, Liquor Ferri Chloroxydi, Liquor Ferri Dialysatus, Liquor Ferri Oxychlorati, Mistura Ferri Amara, Mistura Ferri Aromatica, Mistura Chalybeata, Mistura Ferri cum Magnesi Sulphate, Mistura Ferri Subchloridi, Tinctura Ferri Chlorati Ætherea, and Tinctura Ferri Muratis

Foreign Pharmacopœias—Official in Austr, sp gr 1 280 to 1 290 (10 p c of Iron), Belg, sp gr 1 28 (29 p c Iron Chloride), Dan and Swed, sp gr 1 298 to 1 302 (10 p c of Iron), Dutch, sp gr 1 470 to 1 482 (75 p c of Iron Chloride), Fr, Port and Span, sp gr 1 260 (about 9 p c of Iron), Jap and Norw, sp gr 1 280 to 1 282 (10 p c of iron), Mex, sp gr 1 260 (26 p c of anhydrous Iron Chloride), Swiss, sp gr 1 280 to 1 290 (about 10 p c of Iron), Ger, Hung and Russ sp gr 1 280 to 1 282 (10 p c of Iron), Ital, sp gr 1 29 (10 p c of Iron, 29 p c Iron Chloride), U S, sp gr 1 315 at 25° C (10 p c of Iron, 29 p c Iron Chloride)

Tests—Strong Solution of Ferric Chloride has, *BP*, a specific gravity of about 1 42, but there is a difference between the Pharmacopœia figure for Oxide and this. The *USP* Liquor has a specific gravity of 1 280 to 1 290 at 25° C (77° F), the *PG* Liquor 1 280 to 1 282. The diluted solution answers the tests distinctive of Ferric salts given under Ferrum. Silver Nitrate Solution produces in the diluted solution, acidified with Nitric Acid, a white curdy precipitate, insoluble in Nitric Acid, but readily soluble in Ammonia Solution. It is officially required to indicate 32 0 p c w/v of Iron Oxide, which is equivalent to 39 8 p c of anhydrous Ferric Chloride. The *USP* preparation is required to contain not less than 29 p c of Ferric Chloride, corresponding to 10 p c of metallic Iron. The method of determination adopted by the *BP* is a gravimetric one. The Iron is precipitated as Hydroxide, ignited and weighed as Oxide, the precipitate obtained by adding an excess of Ammonia Solution to a measured quantity of 5 c c diluted with 80 c c of Water, when well washed and dried should leave a residue weighing 1 6 grammes. The *USP* process is volumetric, and is described below. The amount of Oxide yielded by the *BP* process has been shown (*CD* '99, ii 220, '00, ii 163, *YBP*, '99, 361, *PJ*, '99, ii 44, 63, 133, '00, ii 106) to be at variance with the specific gravity. A sample made strictly in accordance with the *BP* gave 1 604 grammes of Iron Oxide per 5 c c, but possessed a specific gravity of 1 480 instead of 1 42.

The more generally occurring impurities are Ammonium, Arsenic, Calcium, Copper, Lead, Potassium, Sodium and Zinc, Nitrates and Ferrous salts. Of these the more important are Arsenic, Copper, Lead and Zinc. Arsenic may be detected by Bettendorf's test with Stannous Chloride Solution, Copper and Zinc by Hydrogen Sulphide, using the filtrate after removal of the Iron by Ammonia Solution, or by Potassium Ferrocyanide Solution after removal of the Iron with excess of Ammonia Solution, Lead by precipitation as Sulphate.

Nitrates are examined for by the Ferrous Sulphate test, and Ferrous salts with Potassium Ferrocyanide Solution.

A test for Oxychloride with Tenth-normal Volumetric Sodium Thiosulphate Solution is given in the *PG* and *USP*. Each of these tests is given in small type below.

Zinc Iodide Starch Solution—Strips of paper soaked with Zinc Iodide Starch Solution should not be coloured blue when brought near to Liquid Ferric Sesquichloride, *PG*.

Potassium Ferrocyanide—Diluted Solution of Ferric Chloride yields a dark blue precipitate with T.S. of Potassium Ferrocyanide, *PG* and *USP*. If 5 c.c. of Ferric Chloride Solution diluted with 20 c.c. of Water be mixed with excess of T.S. of Ammonia and the mixture filtered a colourless filtrate should be obtained, which when supersaturated with Acetic Acid should not be affected by T.S. of Potassium Ferrocyanide, *PG*.

Potassium Ferricyanide—Solution of Ferric Chloride diluted with 10 parts of Water and acidulated with Hydrochloric Acid should not produce a blue coloration with T.S. of Potassium Ferricyanide, *PG*. The *USP* directs that a few drops of freshly prepared T.S. of Potassium Ferricyanide be added to a diluted portion of Solution of Ferric Chloride (1-20), a pure brown colour should be produced which should not turn green or greenish blue at once—*USP*.

Sodium Thiosulphate—Three drops of Ferric Chloride Solution heated slowly to boiling with 10 c.c. of Tenth normal Volumetric Solution of Sodium Thiosulphate should not give a precipitate of Ferric Hydroxide, *USP* and *PG*.

Residue—If to Ferric Chloride Solution (5 c.c. diluted with 20 c.c. of Water, *PG*) there be added excess of T.S. of Ammonia, and the mixture filtered, a colourless filtrate should be obtained which should not yield a weighable residue on evaporation and gentle ignition, *USP* and *PG*.

Hydrogen Sulphide—A portion of the filtrate obtained as described in the Residue test above should not yield a precipitate with T.S. of Hydrogen Sulphide, *USP*.

Barium Nitrate—Another portion of this filtrate should not be affected by T.S. of Barium Nitrate, *PG*.

Ferrous Sulphate—A third portion of 2 c.c. of this filtrate, mixed with 2 c.c. of Sulphuric Acid and 1 c.c. of T.S. of Ferrous Sulphate carefully poured over this as a layer, should not give a brown ring, *PG*. The *USP* directs that a clear crystal of Ferrous Sulphate be added to a cooled mixture of equal volumes of Sulphuric Acid and a diluted portion of Solution of Ferric Chloride Solution (1-10), the crystal should not become coloured brown, nor should a brownish-black colour develop around it.

Stannous Chloride—A mixture of 1 c.c. of Ferric Chloride Solution and 3 c.c. of T.S. of Stannous Chloride should not assume a dark colour in the course of an hour, *PG*.

Volumetric Determination—The *USP* gives the following instructions: 10 grammes of the Solution are diluted to measure 100 c.c. and 11.1 c.c. of this mixture are introduced into a glass stoppered bottle of 100 c.c. capacity, together with 10 c.c. of Water and 2 c.c. of Hydrochloric Acid. 1 gramme of

Potassium Iodide is then added and the mixture kept at a temperature of 40°C (104°F) for half an hour, then cooled and mixed with a few drops of T S of Starch. The mixture when titrated with Tenth-normal Volumetric Solution of Sodium Thiosulphate should require not less than 20 c c of the Volumetric Solution to discharge the blue or greenish colour, 1 c c of the V S is equivalent to 0.5 p c of metallic Iodine.

Preparations

LIQUOR FERRI PERCHLORIDI. SOLUTION OF FERRIC CHLORIDE

Dilute 1 of Strong Solution of Ferric Chloride with Distilled Water to make 4 of a liquid, sp gr 1.110 (1 in 4)

Dose—5 to 15 minims = 0.32 to 1.0 gramme

This solution and the 'Tincture of Ferric Chloride' contain identical proportions of Ferric Chloride, for 'Prescribing Notes' see below

TINCTURA FERRI PERCHLORIDI.—TINCTURE OF FERRIC CHLORIDE *NO Syn*—STEEL DROPS TINCTURE OF STEEL

Mix 1 of Strong Solution of Ferric Chloride with 1 of Alcohol (90 p c), and add Distilled Water to make 4 (1 in 4)

Medicinal Properties.—Astringent, tonic, hæmostatic. Given in passive hæmorrhage and to arrest hæmorrhage in typhoid. As a general tonic during convalescence, highly useful in anæmia, valuable in large doses for faecal and for erysipelatous inflammations. A rectal injection of 60 minims of the Tincture in half a pint of Water kills thread-worms.

Recommended (*L* '04, 11 1178, 1248, 1415), not only in puerperal septicæmia, but also in local and general septic infection occurring in gynecological practice. 15 to 25 minims every two hours strongly recommended in blood poisoning.—*L* '04, 11 1318.

If Potassium Iodide be added to an aqueous solution of Potassium Citrate, and Tincture of Ferric Chloride added, a yellowish-green solution is obtained, containing no free Iodine, and remaining permanent for months at least. This is suggested (*CD* '05, 11 971, *PJ* '05, 11 861) as a means of overcoming the incompatibility of Potassium Iodide and Ferric Chloride solutions.

Dose—5 to 15 minims = 0.3 to 0.9 c c

Prescribing Notes.—Preparations of Iodine can be given in Infusion of Quassia, or Calumba, but they tinge Infusion of Chiretta and Hops, and change to brown or black those of Cusparia, Gentian, Orange, Cascarella, Cinchona, Cloves, Digitalis, and all astringent infusions.

Glycerin is better than an equal quantity of Syrup for masking the unpleasant astringent taste of Ferric Chloride Solutions. Chloroform Water is also useful.

Equal volumes of Liquor Ferri Perchloridi and Glycerin forms a good paint in faecal inflammation.

Styptic Wool, containing Ferric Chloride, is useful for local application.

Incompatibles.—Alkalies and their Carbonates, Lime Water, Calcium Carbonate, Magnesia, Salicylates, Mucilage of Acacia.

Foreign Pharmacopœias.—Official in Dan, Norw and Swed (Solutio Chloreti Ferrici Spirituosa), Dan and Swed, also (Solutio Chloreti Ferrici Spirituoso-Ætherea), Ger and Russ (Tinct Ferri Chlorati Ætherea), US (Tinctura Ferri Chloridi), Port, from the salt, with Alcohol and Ether, Ital (Soluzione Alcolico-Ætherea di Cloruro Ferrico), and in the Soluor with Alcohol and Ether, Swiss (Spiritus Æthereus Ferri). None in the others. See also 'Tinctura Ferri Chlorati Ætherea'.

Tests—Tincture of Ferric Chloride has a specific gravity of 1.085 to 1.089, contains about 12 p c w/v of total solids and about 22 p c w/v of Absolute Alcohol. It yields with Ammonia Solution a reddish brown voluminous precipitate, with Potassium Ferrocyanide Solution a blue precipitate, and with Silver Nitrate Solution a white curdy precipitate, insoluble in Nitric Acid. The *USP* Tincture has a specific gravity of about 1.005 at 25° C (77° F), and contains not less than 13.28 p c of anhydrous Ferric Chloride, corresponding to 4.6 p c of metallic Iron.

Tinctura Ferri Sesquichloridi PL—Tinctura Ferri Murialis P.E.—There is an idea, which periodically finds its way into print, that a Tincture made according to the formula of the London and Edinburgh Pharmacopœias is more efficacious than the *BP*, and can be given in cases where the other is not tolerated. From a chemical point of view the only difference is that *PL* is three-fourths the strength of *BP*, and when freshly made contains one-fifteenth of the Iron in the Ferrous condition. Alcohol has no reducing action on Ferric Chloride, even after years of contact.

Liquor Ferri Chloroxydi and **Liquor Ferri Dialysatus** have been much used as palatable, non-astringent, and non-irritant hæmatics, given in cases where the astringent salts would derange the stomach.

Not Official

LIQUOR FERRI CHLOROXYDI—A solution in Water of a basic Ferric Chloride, containing 0.8 p c of Chlorine for 5 p c of Ferric Oxide, approximating to the formula $\text{Fe}_2\text{Cl}_6 \cdot 7\text{Fe}_2\text{O}_3$. This is the ratio of the Solution made by us many years previous to the use of 'Dialysed Iron'. It was and is still made to contain 7.1 p c of Ferric Oxide to correspond with the official Tincture.

Dose—10 to 30 minims = 0.6 to 1.2 c c

LIQUOR FERRI DIALYSATUS (Dialysed Iron)—This was formerly official in *BP*, but is now omitted. It contains 5 p c of Ferric Oxide, and was dialysed until nearly tasteless. It is better to work to a definite percentage of Chlorine, it may be reduced to 0.3 p c without interfering with the stability of the solution. It is very doubtful, however, whether there is any advantage in reducing the Chlorine ratio below that of **Liquor Ferri Chloroxydi** as described above.

Another method is to add a certain proportion of diluted Ammonia Solution to a solution of Ferric Chloride, so that the precipitate which first forms just redissolves. The Ammonia becomes Ammonium Chloride and the Iron a very basic Oxychloride, from which the Ammonium salt is readily dialysed. Where a saving of expense is an object, as in some large institutions, it would probably be equally efficacious without dialysis.

Dose—10 to 30 minims = 0.6 to 1.8 c c

Foreign Pharmacopœias—Official in Austr (*Ferrum Hydroxydatum Dialysatum Liquidum*); Ger, Hung, Russ and Swed, when **Liquor Ferri Oxidati Dialysati** is prescribed, **Liquor Ferri Oxychlorati** (sp gr 1.050) may be dispensed, Belg and Swiss (*Ferrum Oxychloratum Solution*), sp gr 1.05, Mex (*Oxido de Hierro Dialysado*), sp gr 1.046. Not in the others.

Liquor Ferri Oxychlorati—Dilute Ferric Chloride Solution 85, with 160 of Distilled Water, and pour into a mixture of Ammonia Water 85 and Distilled Water 920, wash, press, and dissolve the precipitate in 8 of Hydrochloric Acid, finally warming it to about 40° C and dilute the solution with Distilled Water until it has a sp gr 1.05. *Ger* *Jap* has the same formula, but employs 2.5 of Hydrochloric acid (80 p c) in place of the *PG* (25 p c).

Liquor Ferri Oxychloridi—Solution of Ferric Chloride (*USP*), 85, by weight, Ammonia Water (*USP*), 85, by weight, Hydrochloric Acid (*USP*), 2.85, by weight, Water, *qs* to produce 100 by weight—*USNF*.

Strong Solution of Ferric Chloride (*B P*), by weight, 22 50, Solution of Ammonia (*B P*), by weight, 35 00, Hydrochloric Acid (*B P*), by weight, 2 35, Distilled Water, *q s* to produce by weight 100—*B P C*

Both of the above are stated to correspond in strength with **Liquor Ferri Oxychlorati**—*P G*

GLYCERINUM FERRI PERCHLORIDI—Solution of Ferric Chloride, 1 fl oz, Glycerin, 1 fl oz—*Middlesex and University*
Ferric Chloride, 1, Glycerin, 4—*Guy's*

MISTURA FERRI AROMATICA—Solution of Ferric Chloride, 10 minims, Aromatic Spirit of Ammonia, 20 minims, Syrup, 40 minims, Water, to 1 fl oz—*St Thomas's*

Dose—1 fl oz Mix with the Syrup the Iron Solution, and add the Aromatic Spirit previously diluted with the Water

This has been incorporated in the *B P C* under the title **Mistura Ferri Ammoniata**

MISTURA CHALYBEATA—Solution of Ferric Chloride, 15 minims, Syrup, 80 minims, Infusion of Quassia, to 1 fl oz—*St Thomas's*

Dose—1 fl oz

This has been incorporated in the *B P C* under the title **Mistura Ferri Amara** with the *syn* **Mistura Chalybeata**

Mistura Ferri Amara—Solution of Perchloride of Iron, 20 minims, Spirit of Chloroform, 5 minims, Infusion of Quassia, to 1 fl oz—*Loch*

MISTURA FERRI CUM MAGNESII SULPHATE—Solution of Ferric Chloride, 15 minims, Magnesium Sulphate, 20 grains, Glycerin, 40 minims, Infusion of Quassia, to 1 fl oz—*St Thomas's*

Dose—1 fl oz

This has been incorporated in the *B P C*

MISTURA FERRI SALINA—Potassium Citrate, 22 grains, Solution of Ferric Chloride, 24 minims, Chloroform Water, to 1 fl oz—*University*

Dose— $\frac{1}{2}$ to 1 fl oz

SYRUPUS FERRI SUBCHLORIDI—Iron Wire, 300 grains, Hydrochloric Acid, 2 fl oz, Citric Acid, 10 grains, Distilled Water, 10 fl drms, Syrup, *q s* to produce 20 fl oz—*B P '85*

Dose— $\frac{1}{2}$ to 1 fl drms = 1 8 to 3 6 cc

This has been incorporated in the *B P C*

TINCTURA FERRI CHLORATI ÆTHEREA—Iron Chloride Solution, 1, Ether, 2, Alcohol (90 p c), 7 All by weight—*Ger, Ital and Jap*

FERRI PERNITRATIS LIQUOR.

SOLUTION OF FERRIC NITRATE

A reddish-brown liquid, readily miscible with Water It contains Ferric Nitrate, $\text{Fe}_2\text{6NO}_3$, eq 480 68, in solution

Medicinal Properties.—Tonic, astringent and escharotic Like the Ferric Chloride it is useful in hæmatemesis and in 'pyrosis' from the bowel, either by the mouth or as an injection with starch mucilage

Dose.—5 to 15 minims = 0 3 to 0 9 cc

110 minims contain $3\frac{1}{2}$ grains of Iron, 100 cc. contain 3 3 grammes

Foreign Pharmacopœias —Not in any

Tests —Ferric Nitrate Solution has a specific gravity of 1.107 to 1.109. It should answer the tests distinctive of Ferric salts given under Ferrum. The diluted solution, when mixed with an equal volume of Sulphuric Acid, keeping the liquid cool meanwhile, yields a dark-brown ring when a solution of Ferrous Sulphate is gently floated on to the surface of the liquids. It is officially required to yield 4.6 p.c. w/v of Iron Oxide, when a measured quantity of 5 c.c. is precipitated with Ammonia Solution in excess, and the precipitate is washed, dried, ignited, and weighed, the residue amounting to 0.23 gramme. It should be free from the impurities mentioned under Liquor Ferri Perchloridi Fortis, Nitrates excepted.

FERRI PHOSPHAS.

IRON PHOSPHATE

A dull, greyish-blue, amorphous, odourless powder, which is officially required to contain not less than 47 p.c. of Hydrrous Ferrous Phosphate, $\text{Fe}_2(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$, eq. 498.48 together with Ferric Phosphate and Iron Oxide.

Solubility —Insoluble in Water, but soluble in Hydrochloric Acid.

Medicinal Properties —A valuable hæmatinic tonic. Given in anæmia, in amenorrhœa, some forms of dyspepsia, rachitis and tubercular bone diseases, in nervous depression and exhaustion with tendency to phosphaturia, and during convalescence.

Dose —5 to 10 grains = 0.32 to 0.65 gramme.

Prescribing Notes —*Given in cachets, pills, or powders. A good pill can be made by adding one third of its weight of 'Diluted Glucose'.*

Official Preparations —Syrupus Ferri Phosphatis, Syrupus Ferri Phosphatis cum Quinina et Strychnina.

Not Official. —Syrupus Triplex, Syrupus Tres, Elixir Ferri Quinina et Strychnina Phosphatum, Glyceritum Ferri Quinina et Strychnina Phosphatum, Pilula Ferri Quinina et Strychnina Phosphatum, Liquor Ferri Phosphatis Fortis, Pilula Trium Phosphatum, Syrupus Ferri Phosphatis Compositus, Squire's Chemical Food, Syrupus Ferri Phosphatis c. Manganeseo Ferri Phosphas Solubilis, Ferri Pyrophosphas Solubilis.

Foreign Pharmacopœias —Official in Spain (Fosfato de Hierro), U.S. (Soluble Ferric Phosphate), Mex. (Fosfato Ferrico - Ferrico). Not in the others.

Tests —Iron Phosphate dissolves in Hydrochloric Acid, yielding a solution which gives with Potassium Ferrocyanide and with Potassium Ferricyanide Solutions the tests distinctive of Ferric and Ferrous salts given under Ferrum. On the addition of Tartaric Acid and an excess of Ammonia Solution it yields on the subsequent addition of Magnesium Ammonio-sulphate Solution a white granular precipitate. It is officially required to contain not less than 47 p.c. of hydrrous Ferrous Phosphate, as determined by the titration of a solution of 1

gramme in Hydrochloric Acid, with Volumetric Potassium Bichromate Solution, using Potassium Ferricyanide Solution as an indicator, at least 28.2 cc of the Volumetric Solution should be necessary. Arsenic may be present as an impurity, and may be tested for by the Bettendorff's test.

Preparations

SYRUPUS FERRI PHOSPHATIS. SYRUP OF FERROUS PHOSPHATE

Iron, in Wire, 75 grains, Concentrated Phosphoric Acid, 1½ fl oz, Syrup, 14 fl oz, Distilled Water, q s to make 20 fl oz of a pale green syrupy liquid, containing 1 grain of anhydrous Ferrous Phosphate in 60 minims.

Dose —½ to 1 fl dm = 18 to 36 cc

This Syrup can be conveniently made by adding 1 volume of Liquor Ferri Phosphatis Fortis to 5½ vols of Simple Syrup and 1½ vols of Distilled Water.

Ferrous Phosphate absorbs Oxygen with great rapidity on exposure to air, and requires such a large excess of Acid to keep it in solution that in framing a formula for Syrupus Ferri Phosphatis a compromise must be made between liability to deposit on the one hand and acidity on the other. We think it is better to use a comparatively small excess, and keep the Syrup in small bottles lying down.

SYRUPUS FERRI PHOSPHATIS CUM QUININA ET STRYCHNINA. SYRUP OF PHOSPHATE OF IRON WITH QUININE AND STRYCHNINE

Iron, in Wire, 75 grains, Concentrated Phosphoric Acid, 1½ fl oz, Quinine Sulphate, 130 grains, Syrup, 14 fl oz, Distilled Water, q s to make 20 fl oz of a pale yellowish-green syrupy liquid, possessing a very bitter taste, and having a strong fluorescence, it contains 1 grain of anhydrous Ferrous Phosphate, ½ grain of Quinine Sulphate, and ⅓ grain of Strychnine in 60 minims.

Dose —½ to 1 fl dm = 18 to 36 cc

It resembles the compound known as Easton's Syrup.

It can be made extemporaneously by dissolving 2 grains of Strychnine and 51 grains of Quinine Sulphate in 24 minims of Concentrated Phosphoric Acid and Distilled Water to 1½ fl oz, Liquor Ferri Phosphatis Fortis, 1 fl oz, Syrup, to make 8 fl oz.

This formula has been incorporated in the *BPC* under the title *Liquor Quininæ et Strychninæ*.

Foreign Pharmacopœias —Official in U.S. Not in the others.

A mixture of Easton's, Fellows' and Parrish's Syrups is sold as 'Triple Syrup'.

Syrupus Triplex —Syrupus Ferri Phosphatis Compositus, 2 fl oz, Syrupus Ferri T. 1 fl oz, Syrupus Eastoni, 1 fl oz —*Pharm. Form.* The Syrupi Tres of the *Edinburgh Royal Infirmary* is the same.

Martindale (1904) and *BPC* (1907) give the proportions as an equal volume of each.

Not Official.

ELIXIR FERRI QUININÆ ET STRYCHNINÆ PHOSPHATUM. —Soluble Ferric Phosphate, 1.75, Strychnine, 0.0275, Phosphoric Acid, 0.20, Ammonium Carbonate, 0.90, Alcohol (95 p.c.), 6.00; Acetic Acid,

2 865, Ammonia Water, *q s*, Distilled Water, *q s*, Aromatic Elixir, *q s* of each to produce 100—*U S P*

Average Dose—1 fl dr̄m = 3 6 c c

This has been incorporated in the *B P C* with slight modification

Soluble Iron Phosphate, 1 75, Quinine, 0 875, Strychnine, 0 0285, Concentrated Phosphoric Acid, 0 25, Ammonium Carbonate, 0 90, Alcohol, 6 25, Acetic Acid, 2 75 Solution of Ammonia, Distilled Water and Aromatic Elixir, *q s* of each to produce 100—*B P C*

In the *B P C Supplement* the 1 75 of Soluble Iron Phosphate is replaced by 1 35 of Ferric Citrate, and sufficient Sodium Phosphate to give the solution a distinctly green coloration

GLYCERITUM FERRI QUININÆ ET STRYCHNINÆ PHOSPHATUM (*U S*)—Soluble Ferric Phosphate, 8, Quinine, 10 4, Strychnine, 0 08, Phosphoric Acid, 20, Glycerin, 50, Water, *q s* to make 100—*U S P*

The **Average Dose**, 15 minims, contains about 1½ grains of Ferric Phosphate, 1½ grains of Quinine, ⅛ grain of Strychnine

This has been incorporated in the *B P C*

It is used for preparing Syrupus Ferri Quininae et Strychninae Phosphatum, *U S P*, by mixing Glycerite 1 with Syrup 3, but the Syrup so produced is very different from the corresponding preparation in the *B P* It contains in each fl dr̄m about 1½ grains of Ferric Phosphate, 1½ grains of Quinine, ⅛ grain of Strychnine

LIQUOR FERRI PHOSPHATIS FORTIS—Containing 8 grains per fl dr̄m of the Anhydrous Phosphate, is made by dissolving 360 grains of Iron Wire in 6 fl oz of Concentrated Phosphoric Acid, with sufficient Water to make 12 fl oz

PILULA FERRI QUININÆ ET STRYCHNINÆ PHOSPHATUM—Ferrous Phosphate, 1 grain, Quinine Sulphate, 1 grain, Strychnine, ⅛ grain, Milk Sugar, 1½ grains, Concentrated Phosphoric Acid, *q s* for one pill—*Martindale*

It is also made half this strength, and either may be combined with Arsenious Acid ⅛ grain—*Martindale*

Pilulae Ferri Phosphatis cum Quinina et Strychnina. *Syn.* **EASTON'S PILLS**—In 100 parts, Ferrous Phosphate, 20, Quinine Sulphate, 20, Strychnine, 0 62, Milk Sugar, a sufficient quantity, Concentrated Phosphoric Acid, *q s* to form a mass Divide into pills containing 2 grains each—*B P C*

Note—Easton's Pills are sometimes made twice the size specified above, and they may be ordered with the addition of ⅛ grain of Arsenious Acid—*B P C*

Although the *B P C* form resembles that of *Martindale*, it is not nearly so definite in strength

Pilula Trium Phosphatum **EASTON'S PILL**—Iron Phosphate, 1 grain, Quinine Sulphate, 1 grain, Strychnine, ⅛ grain, Concentrated Phosphoric Acid, 1½ minims, Liquorice Powder, to 4 grains—*Guy's*

SYRUPUS FERRI PHOSPHATIS COMPOSITUS—Iron Wire, free from Oxide, 87½ grains, Concentrated Phosphoric Acid (sp gr 1 5), 1 fl oz, Distilled Water, 5 fl dr̄m, dissolve by a gentle heat in a flask plugged with Cotton-Wool, the Iron being completely covered by the liquid

Precipitated Calcium Carbonate, 120 grains, Concentrated Phosphoric Acid, 4 fl dr̄m, Distilled Water, 2 fl oz, mix, and add Potassium Bicarbonate, 9 grains, Sodium Phosphate, 9 grains, filter, and set aside

Cochineal, 30 grains, Distilled Water, 7½ fl oz, boil for 15 minutes, and when cooled filter, pouring over the filter a sufficient quantity of Distilled Water to produce 7 fl oz of filtrate to this add Refined Sugar, 14 oz, heat till dissolved, and strain When cold, add the Iron and Calcium Solutions and sufficient Distilled Water to produce 20 fl oz—*B P C Formulary* 1901

This has been incorporated in the *B P C*, employing 288 minims of Orange Flower Water in place of that quantity of Distilled Water in the quantities given above, but the *B P C Supplement* has since altered the quantity to 480 minims

Each fl drm = $\frac{1}{2}$ grain Ferrous Phosphate and $\frac{1}{4}$ grain Calcium Phosphate with small quantities of Potassium and Sodium Phosphates. It should be kept in bottles quite full.

Dose — $\frac{1}{2}$ to 2 fl drm = 18 to 71 cc

SYRUPUS FERRI PHOSPHATIS COMPOSITUS, SQUIRE (Squire's Chemical Food) — The preparation, made for many years by Squire, was imported and subsequently purchased by Squire.

It contains Ferrous Phosphate, Calcium Phosphate, Sodium Phosphate and Potassium Phosphate.

Dose — $\frac{1}{2}$ to 1 teaspoonful, in Water, with meals.

A formula was published many years ago, but how far this has been a success is shown by comparing the Syrups commercially sold, all of them more or less empirically stated to be made according to the published formula.

In nine samples analysed, the Iron Phosphate ranged from 0.19 to 0.66, the Calcium Phosphate from 0.5 to 1.6, the total Phosphoric Acid from 1.5 to 4.7, these results are expressed in grains per fl drm.

Medicinal Properties — A tonic in debility, of whatever origin, and during convalescence from acute diseases. Specially indicated in tuberculosis and rickets, and during pregnancy.

SYRUPUS FERRI PHOSPHATIS C MANGANESIO — Dissolve 100 grains Manganese Phosphate in $1\frac{1}{2}$ fl oz of Liquor Ferri Phosphatis Fortis and 80 minims of Phosphoric Acid, then dilute to 20 fl oz with Simple Syrup.

This Syrup will contain in each fl drm $\frac{1}{2}$ grain each of anhydrous Ferrous Phosphate and anhydrous Manganese Phosphate.

Dose — 1 fl drm = 86 cc

This can sometimes be taken when Syrup of Ferrous Phosphate disagrees.

FERRI PHOSPHAS SOLUBILIS Soluble Ferric Phosphate (*U S*) — Prepared by dissolving Ferric Citrate and Sodium Phosphate in Distilled Water, evaporation and scaling on plates of glass. The scales are transparent and of a bright green colour, freely soluble in Water, they, however, become dark and discoloured on exposure to air. It is used in the preparation of Elixir Ferri Quininae et Strychninae Phosphatum (*U S*), Glyceritum Ferri Quininae et Strychninae Phosphatum (*U S*), and Syrupus Ferri Quininae et Strychninae Phosphatum (*U S*) (from Glycerite).

FERRI PYROPHOSPHAS SOLUBILIS Soluble Ferric Pyrophosphate (*U S*) — Prepared by dissolving Ferric Citrate and Sodium Pyrophosphate in Distilled Water, evaporation and scaling on plates of glass. The scales are transparent and of an apple green colour, freely soluble in Water.

FERRI SULPHAS.

FERROUS SULPHATE

FER, SULFATE DE PROTOXIDE DE FER OFFICINAL, GER, FERROSULFAT, ITAL, SOLFATO FERROSO, SPAN, SULFATO FERROSO

$\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, eq 276.10

Large, translucent, pale green, octonuclear monoclinic prisms, having a saline, styptic, ferruginous taste.

Ferrum Sulphuricum Præcipitatum (*Austri*), resembles the Ferri Sulphas Granulata of *BP '85*, omitted in 1898. It is obtained by pouring an aqueous solution of Ferrous Sulphate into Alcohol (90 p c).

Solubility. — 1 in $1\frac{1}{2}$ of Water, the solution rapidly oxidises on exposure, insoluble in Absolute Alcohol or Alcohol (60 p c), hence it cannot be dissolved in Tinctures.

Medicinal Properties —A powerful astringent and a hæmâtinic tonic, but is apt to irritate the stomach. Internally it is given in anæmia, amenorrhœa, and general debility, along with Quinine it promotes the appetite, given with cathartics, such as Magnes Sulph and Aloes, to increase their action, but at the same time reduce their dose, externally it is used as a lotion for ulceration and erysipelatous surfaces, 3 to 5 grains in an oz of Water, also as an injection for urethral and vaginal inflammations and prolapse of rectum.

Dose —1 to 5 grains = 0.06 to 0.32 gramme

Prescribing Notes —*Given in solution or more generally pill form, to avoid gastric irritation. The Dried Sulphate is best in pills, 8 grains, which are equal to 5 of the crystallised salt, make a nice pill with 'Diluted Glucose'.*

Liquor Ferri Persulphatis is an excellent styptic. 2 grains of Ferrous Sulphate, 30 grains of Magnesium Sulphate, 5 minims of Diluted Sulphuric Acid, Chloroform Water or Peppermint Water to 1 oz, occurs in Hospital formulas as Mistura Ferri Aperiens.

Official Preparations —Ferri Sulphas Exsiccatus and Liquor Ferri Persulphatis. See also 'Ferrum'.

Not Official —Liquor Ferri Subsulphatis (Monsel's Solution), Mistura Ferri et Magnesi Sulphatis, Mistura Ferri Aperiens, Monsel's Salt, Gossypium Ferratum, and Ferri et Ammonii Sulphas.

Foreign Pharmacopœias —Official in Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Norw, Port, Russ, Span, Swed, Swiss and U.S., Mex (Sulfato Ferroso), Fr, Ger, Jap, Russ and Swiss have a Crude Sulphate, Austr has a Precipitated Sulphate, U.S. a Granulated Sulphate.

Tests —Ferrous Sulphate should dissolve to form a clear solution in less than 2 parts of cold Water. The aqueous solution answers the tests distinctive of Ferrous salts given under the heading of Ferrum. Acidified with Hydrochloric Acid its solution yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid.

It is officially required to contain 99.4 p.c. of crystallised Ferrous Sulphate as determined by titration with Volumetric Potassium Bichromate Solution, using Potassium Ferricyanide Solution as an indicator. The *U.S.P.* salt is required to contain not less than 99.5 p.c. pure Ferrous Sulphate. The processes of determination are compared in the small type below under the heading of Volumetric Determination.

The more generally occurring impurities are Ammonium, Copper, Potassium, Sodium and Zinc, and Ferric salts. They are best detected by oxidising the aqueous solution with Nitric Acid and precipitating the Iron as Ferric Hydroxide with Ammonia Solution. The filtrate should not be blue in colour nor yield a precipitate on the addition of Hydrogen Sulphide, indicating the absence of Copper and Zinc, another portion of the filtrate evaporated to dryness and ignited should not leave a weighable residue, indicating the absence of Potassium and Sodium salts. The *B.P.* uses Hydrogen Sulphide as a test for Ferric salts. It should dissolve 1 in 1½ of Water as stated above, and the solubility of the sample in less than 2 parts of cold Water is officially included as a test for the absence of Oxysulphate.

Hydrogen Sulphide—If 2 grammes of the salt in aqueous solution be added to 10 c.c. of Bromine Water, and excess of T S of Ammonia be added to the mixture, the filtrate from this mixture should be colourless and should not be affected by T S of Hydrogen Sulphide, *P G*. This test is also given in the *USP*. 1 gramme of the salt dissolved in 25 c.c. of Water containing 1 c.c. diluted Sulphuric Acid is heated to boiling and oxidised with Nitric Acid. A slight excess of T S of Ammonia is then added and the mixture filtered. A colourless filtrate is obtained which should not respond to the time-limit test for heavy metals. The colourless filtrate obtained as in the preceding test should not on evaporation and ignition leave a weighable residue, *P G* and *USP*.

Volumetric Determination—An aqueous solution of 1 gramme of Iron Sulphate acidified with Sulphuric Acid requires at least 36 c.c. of Volumetric Solution of Potassium Bichromate, *BP*, the *USP* directs that 1.38 grammes of the salt in uneffloresced crystals be dissolved in 25 c.c. of dilute Sulphuric Acid and the solution titrated with Tenth-normal Volumetric Solution of Potassium Permanganate, not less than 49.75 c.c. of the Volumetric Solution should be necessary to impart a permanent pink colour to the liquid.

Preparations

FERRI SULPHAS EXSICCATUS. EXSICCATED FERROUS SULPHATE. DRIED SULPHATE OF IRON—*BP '85*

Ferrous Sulphate, submitted to a temperature of 100° C (212° F), until it ceases to lose aqueous vapour, reduce to a fine powder and keep in dry, well-stoppered bottles. It should be slowly but completely soluble in Water.

Dose— $\frac{1}{2}$ to 3 grains = 0.032 to 0.20 gramme

3 grains are equal to 5 grains of Ferrous Sulphate

Foreign Pharmacopœias—Official in Dan and Swed, dried at 104° to 122° F (40° to 50° C), Ger, Swiss and U.S., dried at 212° F (100° C), Dutch, Fr; Russ and Span, no temperature given. Not in the others.

Tests—Dried Ferrous Sulphate dissolves slowly but completely in Water, the solution answers the tests distinctive of Ferrous salts given under Ferrum. The 1 in 20 aqueous solution acidified with Hydrochloric Acid yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. It is officially required to contain at least 92.5 p.c. of dried Ferrous Sulphate of the official formula, as determined by titration with Volumetric Potassium Bichromate Solution, using Potassium Ferricyanide Solution as an indicator. The processes of the *BP* and *P G* are compared below under the heading of Volumetric Determination. It should, of course, be free from the impurities mentioned under Ferrous Sulphate, but *BP* does not say so. A standard suggested for Arsenic (*CD '08*, 1796) is 2 parts per million.

Volumetric Determination—A solution of 1 gramme of the salt in water acidified with Sulphuric Acid requires at least 54.6 c.c. of the Volumetric Solution of the Potassium Bichromate. This corresponds to at least 92.5 p.c. of Exsiccated Ferrous Sulphate $\text{FeSO}_4 \cdot \text{H}_2\text{O}$, *BP*, the *P G* gives the following directions—Let 0.2 gramme of the salt be dissolved in 10 c.c. of diluted Sulphuric Acid and the solution mixed with Solution of Potassium Permanganate (5-1000) until a faint reddening occurs. After the colour is destroyed, which may be effected if necessary by a few drops of Alcohol, 2 grammes of Potassium Iodide are added and the mixture is allowed to stand for 1 hour at ordinary temperature in a closed vessel. It is then titrated with Tenth-normal

Volumetric Solution of Sodium Thiosulphate, of which at least 10 c c should be necessary for the combination of the free Iodine, *P G*

LIQUOR FERRI PERSULPHATIS. SOLUTION OF FERRIC SULPHATE

Ferrous Sulphate, 16, Sulphuric Acid, $1\frac{1}{2}$, Nitric Acid, $1\frac{1}{2}$, Distilled Water, *q s* to make 22 of a reddish-brown liquid, sp gr 1.441, miscible with Water and Alcohol (90 p c)

Introduced for making several preparations of Iron which are enumerated under 'Ferrum,' p 504

Foreign Pharmacopœias—Official in Ital, Jap and Swiss, sp gr 1.428 to 1.430, Russ, sp gr 1.426 to 1.430, U S, sp gr 1.430 to 1.450 at 25° C (77° F) Not in the others

Tests—Ferric Sulphate Solution is officially required to possess a specific gravity of 1.441. Its diluted aqueous solution should answer the tests for Ferric salts given under Ferrum. It should yield when diluted with 10 times its volume of Water a brown but no pronounced blue coloration with Potassium Ferricyanide Solution, indicating the absence of more than a trace of Ferrous salt. It should not decolorise Potassium Permanganate Solution. It is officially required to yield 1.04 grammes of Iron Oxide, as gravimetrically determined by precipitation as Hydroxide with Ammonia Solution, washing, drying and incinerating. The *U S P* Liquor is required to contain 36 p c of Normal Ferric Sulphate [$\text{Fe}_2(\text{SO}_4)_3$ eq 397.05], corresponding to not less than 10 p c of metallic Iron. The *U S P* employs a volumetric Iodometric method for the determination of Iron. The respective processes are compared in small type below under the headings Gravimetric and Volumetric Determinations.

The liquor should be free from the impurities mentioned under Ferrous Sulphate.

Potassium Ferricyanide.—A small portion of the solution diluted with about 10 volumes of Water should yield with a few drops of freshly-prepared T S of Potassium Ferricyanide a pure brown colour without a tinge of green or greenish-blue, *U S P*.

Sulphuric Acid.—If 2 volumes of the solution be slowly mixed with 1 volume of concentrated Sulphuric Acid in a beaker, no solid white mass should separate, *U S P*.

Ferrous Sulphate and Sulphuric Acid.—If a crystal of Ferrous Sulphate be added to a diluted portion of the solution (about 1-10), mixed with an equal volume of concentrated Sulphuric Acid and cooled, the crystal should not become brown nor should a brownish-black colour develop around it, *U S P*.

Gravimetric Determination—The reddish brown precipitate produced when a measured quantity of 5 c c of the Liquor diluted with 80 c c of Water is treated with Ammonia Solution in excess, should, when washed, dried, ignited, cooled and weighed amount to 1.04 grammes, *B P*.

Volumetric Determination—A weighed quantity of 1.11 gramme is introduced into a glass stoppered bottle of about 100 c c capacity together with 15 c c of Water and 2 c c of Hydrochloric Acid. 1 gramme of Potassium Iodide is added and the mixture kept at a temperature of 40° C (104° F) for half an hour, then cooled and titrated with Tenth normal Volumetric Sodium Thiosulphate Solution using Starch Solution as an indicator, not less than 20 c c shall be required to discharge the blue or greenish colour of the liquid. 1 c c of Tenth normal Thiosulphate = 0.5 p c w/w metallic Iron, *U S P*.

Not Official.

MISTURA FERRI ET MAGNESII SULPHATIS—Sulphate of Iron, 8 grains, Magnesium Sulphate, 30 grains, Dilute Sulphuric Acid, 5 minims, Distilled Water, to 1 fl oz—*Royal Free*

Ferrous Sulphate, 2 grains, Magnesium Sulphate, 20 grains, Diluted Sulphuric Acid, 10 minims, Water, to 1 fl oz—*King's*

Mistura Ferri Aperiens—Ferrous Sulphate, 2 grains, Magnesium Sulphate, 30 grains, Diluted Sulphuric Acid, 2 minims, Peppermint Water, to 1 fl oz—*University*

LIQUOR FERRI SUBSULPHATIS (U S)—An aqueous solution of basic Ferric Sulphate, corresponding to not less than 13.57 p c of Iron. It is known as **Monsel's Solution**

Monsel's Salt is produced by evaporating and scaling the solution

GOSSYPIUM FERRATUM—Moisten Cotton-Wool with Glycerin, then steep the damp Wool in a solution of Ferrous Sulphate, 1 part to 2 parts of Water, squeeze out as much as possible of the liquid, and, without drying, pack the prepared wool into a bottle furnished with a glass stopper

FERRI ET AMMONII SULPHAS Ammonio Ferric Alum—Iron Alum is an Alum in which Iron takes the place of Aluminium. Pale violet octahedral crystals, which are efflorescent. It should contain 99.5 p c of pure uneffloresced Ferric Ammonium Sulphate

Solubility—Soluble 1 in 3 of Water, insoluble in Alcohol (90 p c)

It is used in bleeding from the kidneys, it arrests the hæmorrhage and the anæmia that accompanies it, it is considered more astringent than Alum

The aqueous solution will, even after filtration, deposit unless slightly acidified with Diluted Sulphuric Acid

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Foreign Pharmacopœias—Official in U S

Tests—Ferric Ammonium Sulphate dissolved in Water yields a blue precipitate with Potassium Ferrocyanide Solution and a brownish-red precipitate with Potassium Hydroxide Solution, followed by the evolution of Ammonia gas on warming. The aqueous solution gives with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. A weighed quantity of 0.555 gramme of the uneffloresced crystals is dissolved in 15 c c of Water and 2 c c of Hydrochloric Acid, in a 100 c c glass stoppered flask, 1 gramme of Potassium Iodide is added, the flask securely fastened, the mixture kept for half an hour at 40° C (104° F), and then cooled, not less than 11.5 c c of Tenth-normal Volumetric Sodium Thiosulphate Solution should be required to discharge the colour. 1 c c of Tenth-normal Thiosulphate = 1 p c of metallic Iron

FERRUM REDACTUM.**REDUCED IRON**

FR, FER REDUIT PAR L'HYDROGÈNE, GER, REDUZIRTES EISEN, ITAL, FERRO RIDOTTO DALL' IDROGENA, SPAN, HIERRO REDUCIDO POR EL HIDROGENO

A fine, tasteless powder, possessing a dull iron-grey metallic appearance, and strongly attracted by a magnet. It is officially required to contain at least 75 p c of metallic Iron, with a variable amount of Iron Oxide. It is prepared by the reduction of Ferric Hydroxide, at a dull red heat, by dry Hydrogen

With reference to the keep of Reduced Iron, it may be noted that, under ordinary atmospheric conditions, a sample containing 91.5 p c of Iron, loosely covered with paper to keep out dust, lost only 1 p c of metallic Iron in a month

Medicinal Properties—Hæmatinic Given in chlorosis and amenorrhœa

As Hydrogen is evolved by its contact with the acid gastric secretion, flatulence may be set up

Dose—1 to 5 grains = 0.065 to 0.32 gramme

Prescribing Notes—*Given in powder, pill, or in lozenges. Pills containing Reduced Iron have a tendency to crack. An excellent pill can be made by mixing Reduced Iron 24 grains, Liquorice Powder 6 grains, Glycerin of Tragacanth 6 grains, and dividing into 12 or more pills as desired.*

Official Preparation—Trochiscus Ferri Redacti

Foreign Pharmacopœias—Official in Austri, Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests—Reduced Iron dissolves in Hydrochloric Acid, and Hydrogen gas is at the same time evolved. The solution thus yielded gives the tests distinctive of Ferrous salts given under Ferrium, and on oxidation with Nitric Acid the tests distinctive of Ferric salts also given under Ferrium.

It is officially required to contain at least 75.48 p.c. of metallic Iron as determined by the titration with Volumetric Potassium Bichromate Solution of the Ferrous Sulphate produced when an excess of Copper Sulphate in solution is decomposed by a weighed quantity of Reduced Iron, Potassium Ferricyanide Solution being used as an indicator. The equivalent amount of Copper is precipitated from a hot solution of 1 gramme of Copper Sulphate in 15 c.c. of Water by the addition of a weighed quantity of 0.25 of a gramme of the Reduced Iron, 10 minutes is allowed for the completion of the reaction. The resulting solution of Ferrous Sulphate is filtered with as little exposure to the air as possible and acidified with Sulphuric Acid, it should require at least 33.7 c.c. of Deci-normal Volumetric Bichromate Solution. Both *USP* and *PG* employ Iodometric methods, which are described below in small type under the heading Volumetric Determination. The *BP* 1898 raised the percentage of metallic Iron from 50 to at least 75.5 p.c., but there is no difficulty in obtaining Reduced Iron containing over 90 p.c. of metal. It has been pointed out (*CD* '99, ii 214, *PJ* '99, ii 109) that the Copper Sulphate method is not satisfactory, and that either one of the other two methods tried is to be preferred (these were the Iodometric process and the 'Mercuric Chloride' process official in the *US* '90, which have now been abandoned in favour of the Iodine method).

The *BP* does not include specific tests for impurities, with the exception of Sulphur, which is officially stated to be recognised by an odour of Hydrogen Sulphide during solution, both *USP* and *PG* employ Lead Acetate paper. The more generally occurring impurities, other than the above, are Arsenic, Copper, Silica, Carbon, and alkali Carbonates. Arsenic may be detected by the modified Gutzzeit's test, after the preliminary precipitation indicated below, or by the Bettendorf's test. It has been suggested that a limit of Arsenic should be included in the *BP*. The *USP* adopts a limit of 1 in 100,000. A limit of 0.05 p.c. has been suggested (*CD* '01, ii 242), and this standard is upheld (*YBP* '03, 242). It is stated (*CD* '08, i 796)

that a standard of 0.02 p.c. appears to be a sufficiently stringent limit for Arsenic in Reduced Iron. Copper may be detected by Hydrogen Sulphide in a solution rendered faintly acid with Hydrochloric Acid, or it may be detected by oxidising the Ferrous salt to the Ferric condition, precipitating the Iron as Ferric Hydroxide with Ammonia Solution and examining the filtrate. The limit might be fixed at 1 in 5,000. Silica and Carbon may be detected by the residue insoluble in Hydrochloric Acid, which should not amount to more than 1 p.c. Alkali salts may be detected by the reaction towards red Litmus paper of the water with which the sample has been shaken, and by the residue left on evaporation of the same after filtration.

Stannous Chloride—Let 0.2 gramme of Reduced Iron and 0.2 gramme of Potassium Chlorate be mixed with 2 c.c. of Hydrochloric Acid in a large test glass, and after the reaction ceases the mixture be warmed until free Chlorine is expelled. The solution is then filtered. 1 c.c. of the filtrate so obtained, with the addition of 3 c.c. of T.S. of Stannous Chloride, should not assume a dark colour in the course of an hour, *P. G.*

Modified Gutzet's Test—The *U.S.P.* gives the following instructions for treating Reduced Iron before proceeding to test for Arsenic.—To 0.5 gramme of Reduced Iron contained in a small covered beaker, add 20 c.c. of diluted Sulphuric Acid. After the reaction has somewhat subsided, warm the liquid on a water-bath until the reaction ceases, then collect any minute undissolved residue of impure Iron Arsenide upon a very small filter, rinse the beaker with Water, add the rinsings to the filter, and wash the residue with Water until free from acid reaction. Transfer the residue to the beaker by rinsing it back, and after adding about 0.25 gramme of Potassium Chlorate and 5 c.c. of Hydrochloric Acid evaporate the solution slowly to dryness on a water-bath. Dissolve the residue in sufficient Water to measure 50 c.c., then add 5 c.c. of this solution to 5 c.c. of a saturated Solution of Sulphurous Acid and heat the liquid on a water-bath for fifteen minutes, until all traces of Sulphurous Acid have been removed. The resulting solution should not respond to the modified Gutzet's test for Arsenic.

Volumetric Determination—A weighed quantity of 0.555 gramme of Reduced Iron is introduced into a 100 c.c. flask containing about 2.6 grammes of Iodine, the weight of which is subsequently accurately recorded, 6 c.c. of Water and 2 grammes of Potassium Iodide are added, the flask is securely stoppered and set aside for one hour, sufficient Distilled Water is added to measure exactly 100 c.c., mixed well, and 25 c.c. of this solution removed, and after the addition of a few drops of Starch Test-solution it is titrated with Tenth-normal Volumetric Sodium Thiosulphate Solution. The weight of Iodine taken is divided by 0.02518, the quotient subtracted from twice the number of c.c. of Tenth-normal Volumetric Sodium Thiosulphate Solution used, and the remainder represents the percentage of Iron in the sample. Instead of employing the official Iodine for the test the percentage of purity of the Iodine may be ascertained by a separate experiment, and the equivalent quantity of pure (100 p.c.) Iodine may be used instead of the 2.6 grammes referred to above, *U.S.P.*

A weighed quantity of 0.3 gramme of finely-powdered Reduced Iron is mixed with 10 c.c. of T.S. of Potassium Iodide, and into this mixture is gradually introduced 1.5 grammes of powdered Iodine, cooling and shaking. As soon as the Iron and Iodine are completely dissolved, the liquid is diluted with Water to 100 c.c., allowed to stand and deposit. A measured quantity of 50 c.c. of the clear Solution is filtrated with Tenth-normal Volumetric Solution of Sodium Thiosulphate, of which not more than 10.3 c.c. should be necessary for combination with the free Iodine, *P. G.*

Preparation

TROCHISCUS FERRI REDACTI. REDUCED IRON LOZENGES

1 grain of Reduced Iron in each, on a Simple Basis

Dose.—1 to 6 lozenges

FERRUM TARTARATUM.**TARTARATED IRON**

FR, FERRITARTRATE DE POTASSIUM, GER., KALIUMFERRITARTRAT, ITAL, TARTRATO FERRICO POTASSICO, SPAN, TARTRATO FERRICO POTASSICO

Thin, deep ruby red, translucent, slightly deliquescent scales, having a sweetish, ferruginous and astringent taste

It should be kept in well closed vessels, of a dark amber tint, and protected as far as possible from air and light

Solubility—1 in 1 of Water (slowly), very sparingly in Alcohol (90 p c)

Medicinal Properties—Chalybeate tonic, and slightly diuretic, suitable in the anæmia of convalescence

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Foreign Pharmacopœias—Official in Belg (Tartras Ferrico-Potassicus), Fr (Ferritartiate de Potassium), Ital (Tartriato Ferrico Potassico), Mex (Tartriato de Potasio y Fierro), Port (Tartriato de Potassa e de Feiro), Russ (Feiro Kalium Tartaricum), Span (Tartrato Ferrico Potassico), US (Ferri et Potassii Tartras) Not in the others

Ferri et Ammonii Tartras is also official in US

Tests—Tartarated Iron dissolves slowly in Water. The solution answers the tests distinctive of Ferric salts given under Ferrum. It is stated to yield no dark blue coloration, but only a greenish turbidity with Potassium Ferricyanide Solution, but it always contains Ferric salt, which precipitates with the Potassium Ferricyanide reagent

If the precipitate produced by Potassium Hydroxide Solution be removed by filtration and the filtrate be slightly acidulated with Acetic Acid, it yields, as it cools, a crystalline deposit, more particularly if the filtrate is first mixed with a little Alcohol (90 p c). It is officially required to yield 30.0 p c of Ferric Oxide, as determined by incinerating a weighed quantity at a red heat, cooling, and washing the residue till free from Potassium Carbonate, which operation is not always an easy matter to complete. The *USP* preparation is required to contain Iron and Potassium Tartrate corresponding in amount to not less than 15 p c of metallic Iron. The *USP* method of determination is an Iodometric one, and is described below. If the Iron be removed from a 1 in 10 aqueous solution by boiling with an excess of Potassium Hydroxide Test-solution, the filtrate, when slightly acidified with Acetic Acid, will gradually deposit a white crystalline precipitate, *USP*

With Potassium Ferrocyanide Test solution, the solution should not afford a blue colour or precipitate, unless it be acidulated with Hydrochloric Acid, *USP*

Ammonia—The aqueous solution of the salt should not yield any precipitate with T.S. of Ammonia, but is rendered darker, *USP*

Volumetric Determination—The *USP* gives the following instructions—If 0.555 gramme of the dry salt be dissolved in 15 c c of Water and 2 c c of Hydrochloric Acid, in a glass stoppered flask having a capacity of about 100 c c, and if after the addition of 1 gramme of Potassium Iodide, and securely

closing the flask, the mixture be kept for half an hour at 40° C (104° F) and then cooled, it should require not less than 15 c c of Tenth-normal Volumetric Solution of Sodium Thiosulphate to discharge the colour of the liquid, Starch T S being used as indicator (each c c of Tenth-normal Volumetric Solution of Sodium Thiosulphate indicating 1 p c of metallic Iron)

FICUS.

FIGS

FR, FIGUL, GER, FEIGEN, ITAL, FICHI, SPAN, HIGOS

The dried fleshy receptacles of *Ficus Carica*, L

Medicinal Properties—Nutritious, laxative, and demulcent
Chiefly used medicinally in constipation Cut open and heated, it forms a convenient cataplasm

Official Preparation—Contained in Confectio Sennæ

Foreign Pharmacopœias—Official in Port (Figs Passados), U S Not in the others

Descriptive Notes.—Dried Figs are usually imported in two forms, natural and pulled figs Natural figs are fruits that have not been made supple by kneading and squeezing, pulled figs are those so treated, and are generally imported in small boxes, in which they have been packed by careful pressure They are flattened, translucent, and have the characteristic taste of the fig In dry weather they are covered with a saccharine efflorescence Greek figs are inferior in quality, containing less pulp, and are smaller

The fruit is sometimes called a syconus, it is a pyriform receptacle filled with female flowers, each of which contains a minute ovary which, when ripe, forms an achene, often erroneously called a seed The male flowers are developed amongst bracts surrounding the minute tubular orifice at the apex

Compound Syrup of Figs—Under this title several preparations are made containing the soluble laxative constituents of Figs and Senna

FILIX MAS.

MALE FERN

The Rhizome of *Aspidium Filix-mas*, Sw, carefully dried

Medicinal Properties.—The powder of the rhizome is slightly tonic and astringent, chiefly used in the form of Liquid Extract as an anthelmintic for tapeworm

Prescribing Notes.—The Liquid Extract, which is an Ointment can be given in Milk, or made into an emulsion with 1 to 2 fl drms of very rich Mucilage of Gum Acacia, or $\frac{1}{2}$ to 1 drms of powdered Acacia, and with Peppermint Water or Milk to form a 2 oz draught, or in capsules Best given in the early morning fasting after a purge on the 11th day, as the worm is so protected by food It is more effective if the dose of 1 drms be continued for 3 or 4 days

and given at intervals of half an hour. It should be followed 12 hours afterwards by a brisk purgative (not Castor Oil) to clear away the dead worm.

Under the headings *Haustus*, and *Mistura*, *Filicis Maris*, several formulas are given in the Pharmacopœias of the London Hospitals.

10 minims of Tincture of Senega recommended for each fl dr̄m of liquid extract, as an emulsifying agent.—*P J '02*, ii 869.

Official Preparation—*Extractum Filicis Liquidum*

Not Official—*Mistura Filicis*

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Jap, Fr (*Fougere*), Ger, Hung, Norw, Ital (*Felce Maschio*) Port (*Feto Macho*), Russ, Mex, Span (*Helecho Macho*), Swed, Swiss, US (*Aspidium*)

Descriptive Notes—The rhizome, as met with in English commerce, is sometimes entire and sometimes cut in half longitudinally for facility in drying, and has the scaly bases of the leaf stalks or stipes attached to it, these being angular and about 1 inch long and about $\frac{1}{2}$ inch in diameter. The official description limits the size from 3 to 6 inches (7.5 to 15 cm) in length, $\frac{3}{4}$ to 1 inch (2 to 2.5 cm) in diameter. The rhizome should be green internally. It should not be kept more than one year. The *BP*, as well as the *PG* and *USP*, requires that the rhizome should not be kept more than a year, as the medicinal activity is decreased on keeping. An indication of its freshness when purchased is the yellowish green tint of the rhizome and stipes when cut transversely. The *BP* requires that it should be collected late in the autumn, and divested of its roots, leaves and dead portions. The *USP* states that the chaff (*ie*, scales), together with the dead portions of the rhizome and stipes, should be removed, and only such portions used as have retained their internal green colour. The distinguishing feature of the rhizome is the presence in the transverse section of 10 large vascular bundles forming a circle, beyond which a number of small ones are scattered, the leaf stalks showing eight only in an irregular circle. In the intercellular spaces near the apex, of the rhizome globular stalked glands are found, which do not occur in most of the allied ferns likely to be mistaken for it in this country, such as *A. Oreopteris*, Sw, and *Athyrium Filix-fœmina*, Roth. The scales of *A. Filix-mas*, Sw, have two glands, at the base, but that of *A. spinulosum*, Sw, which has been found frequently in Germany mixed with the true rhizome, has them also on the margins of the scales. In S Africa the root of *Aspidium athamanticum*, Kunze, is used as a tænicide under the name of 'Unkomokomo,' and in Continental commerce as 'Pannum.'

Tests—The ash of Male Fern varies from 1.5 to 3.0 p c, the insoluble portion of the ash from 0.1 to 0.5 p c. A standard of not less than 5 p c has been suggested.

Preparation

EXTRACTUM FILICIS LIQIDUM LIQUID EXTRACT OF
MALE FERN

Male Fern exhausted by percolation with Ether, and subsequent evaporation of the Ether

It is referred to in the *U S P* as *Oleoresina Aspidii*, and is prepared from *Aspidium* by percolation with Acetone

Dose—45 to 90 minims = 2 7 to 5 4 c c

For larger doses than 90 minims, see *L* '88, 11 1037, *B M J* '89, 1 319, and " as to mode of administration, *L* '94, 11 255

U S states that the granular crystalline substance, which deposits on standing, should be thoroughly mixed with the liquid portion before use

The activity of the Extract is supposed to be due to **Filicic Acid**—*P J* (3) xxii 84, and this varies in different samples from 0 71 to 9 59 p c, reaching in one sample 13 07 p c—*P J* '97, 11 85

Foreign Pharmacopœias—Official in Austri and Russ (Ext *Filicis Maris*), Belg, Dan, Dutch, Ger, Jap, Norw, Swed and Swiss (Ext *Filicis*), *Fr* (Extrait de Fougère Male), Hung (Extract *Filicis Maris Æthereum*), Ital (Estratto di Felce Maschio Etereo), Port (Extracto de Feto Macho Etereo), Span (Aceite de Helecho Macho), *U S* (*Oleoresina Aspidii*) All made with Ether

Test.—Fluid Extract of Male Fern has a specific gravity of 1 008 to 1 010

Not Official

MISTURA FILICIS—Liquid Extract of Male Fern, 1, Powdered Acacia, 1, Chloroform Water, to 8—*University*

FENICULI FRUCTUS.

FENNEL FRUIT

FR, FENOUIL DOUX, *GER*, FENCHEL, *ITAL*, FINOCCHIO,
SPAN, HINOJO

The dried ripe Fruit of *Foeniculum capillaceum*, Gilib, from cultivated plants

Medicinal Properties—Stimulant, aromatic, and carminative
In action similar to Anise Antispasmodic in intestinal colic of children

In infants an infusion (1 to 60) is employed as an enemæ for the expulsion of flatus

Official Preparation—*Aqua Fœniculi* Used in the preparation of *Pulvis Glycyrrhizæ Compositus*

Not Official—*Oleum Fœniculi*

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, *Fr* (Fœnoul Doux), Ger, Hung, Jap, Norw, Ital (Finocchio), Port (Funcho), Russ, Mex and Span (Hinojo), Swed, Swiss and *U S*

Descriptive Notes—A number of different varieties of Fennel fruits are met with in commerce, varying from 3 to 4 mm (*Japaresc*), to 8 to 10 mm (*Saxon*) in length, and from 1 5 mm (*Russian*) to 3 mm (*French and Saxon*) in diameter The larger fruits are usually slightly curved, and greenish or greenish-brown in colour, according to age and ripeness when collected The mericarps are usually united, and taper at the apex, and the longitudinal ridges are prominent with six large vittæ in each mericarp, two being on the flat, and four on the convex side The different kinds vary in the percentage of Oil and in the relative percentage of Anethol and Fenchone that they

contain The Indian fruit, which yields the smallest percentage of oil, is referred to, *F. panmorium*, D C, which, however, is regarded by some botanists as only a variety of *F. capillaceum*

Fennel fruits, after distillation, are used in cattle foods, and to adulterate powdered Fennel fruits The directions given in the *B P* limit the Fennel fruits to those 5 to 10 mm long and 3 mm in diameter, these include the Saxon and the cultivated French varieties, unless the Persian, Indian and Galician, which come within the limits of length, but are rather less in diameter than the official measurement, could be considered to be not excluded by the term 'about' prefixed to the official measurement So far as flavour is concerned the Saxon, French, Macedonian, Persian, and Japanese are the best Under the microscope the powder is characterised by the spiral and reticulated thin-walled colourless parenchymatous cells of the mesocarp, and the obliquely arranged thin walled linear oblong cells, about 3 to 5 times as long as broad, of the inner epidermis, and the absence of striation on the cells of the outer epidermis

Tests—The ash of Fennel Fruit should amount to not more than 10 p c The ash of four samples determined in the author's laboratory amounted to 8.47, 8.93, 9.75 and 7.70 p c The ash of six samples of the Pulvis to 10.85, 12.8, 9.90, 8.91, 13.0 and 9.89 p c Good Fennel fruits yield from 3 to 5 p c of volatile oil

Preparation

AQUA FENICULI FENNEL WATER

Fennel Fruit, 1, Water, 20 Distil 10 (1 in 10)

Dose—1 to 2 fl oz = 28.4 to 56.8 c c

Foreign Pharmacopœias—Official in Austri, 1 in 20, Ital, Mex and Port, 1 in 4, Ger, Jap and Russ, 1 in 30, Hung and Swed, 1 in 10, Dutch and Swiss, 1 in 25, Belg, Oil 1, Alcohol 99, Water 3800, Dan, with Oil, 1 in 2000, and U S, with Oil, 1 in 500 Not in Fin or Norw

Not Official

OLEUM FENICULI—A colourless or slightly yellow liquid, possessing a peculiar characteristic odour and taste It is distilled from Fennel Fruit, the yield of Oil varying from 4 p c to 6 p c

The important constituent of this Oil is Anethol It also contains the ketone, Fenchone, which is isomeric and closely related to Camphor

The commoner oils contain the Terpenes, Dextro pinene and Dipentene, together with Phellandrene and Limonene A good oil contains about 60 p c Anethol

The Oil from Japanese Fennel resembles closely that from the other varieties—*P J* '96, 11 91, *C D* '96, 11 191

Commercial varieties of Fennel and their essential oils—*P J* '97, 1 225

Dose—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Ger, Hung, Jap, Norw, Port, Russ, Swed, Swiss and U S Not in Fr, Ital or Mex

Tests—Fennel Oil has a specific gravity of 0.965 to 0.980 It is soluble in an equal volume of Alcohol (90 p c) and in 10 of Alcohol (80 p c) It is dextrogyrate, the rotation being from -5° to $+20^{\circ}$ in a 100 mm tube. The solidifying point should be between 5° and 10° C (41° and 50° F)

The more generally occurring adulterations of Fennel Oil are oils from which the Stearoptene has been removed, Alcohol, Volatile Oils containing Phenols, and Oil of Turpentine. The addition of oils from which the Anethol has been abstracted is rendered evident by the lowering of the solidifying point, Alcohol by the lowering of the specific gravity, whilst Turpentine is also detected by the reduction in the specific gravity, the Alcohol-solubility and Rotation.

Ferric Chloride T.S. is employed to detect volatile oils containing Phenol, the addition of a drop of the solution to an alcoholic solution of the Oil should produce no coloration.

Not Official

FORMIC ALDEHYDE

METHANAL METHYL ALDEHYDE

Produced by the limited oxidation of Methyl Alcohol. A gas condensable by cold to a clear mobile liquid. The commercial article 'Formol' or 'Formalin' is stated to be a 40 p.c. solution.

The Formaldehyde Solution official in the *U.S.P.* is required to contain not less than 37 p.c. w/w, that of the *P.G.* about 35 p.c. w/w, and that of *Fr. Codex*, 35 p.c. w/w of absolute Formaldehyde.

FORMALDEHYDUM SOLUTUM—A clear colourless fluid, with an irritating odour, containing from 35 to 40 p.c. of Formaldehyde.

Medicinal Properties—The strong solution (35 to 40 p.c.) is a powerful antiseptic, disinfectant and deodorant, it is also a powerful caustic, and should be handled with care. The vapour is irritating to the eyes and nose, probably due to traces of Formic Acid. Even in very dilute solution, 1 of Formic Aldehyde in 20,000, or 1 of Formalin in 8000, it preserves liquids otherwise liable to decay. Mixed with 50 to 100 of Water, may be used as a general antiseptic in the sick room for washing the hands, spray, etc., and with 400 to 500 of Water as an antiseptic mouth-wash or gargle.

Case of poisoning by drinking 4 oz. of a 4 p.c. solution—*P.J.* '99, ii 295.

Formalin (40 p.c.) in 2000 to 3000 of Water used freely to hypopyon ulcers, and septic abrasions of the cornea—*B.M.J.* '96, i 144.

2 p.c. solution in ringworm—*B.M.J.E.* '94, ii 108, *Y.B.T.* '95, 894.

40 p.c. solution applied to ringworm—*B.M.J.* '96, ii 650.

40 p.c. solution sometimes causes suppuration, and is not so useful for ringworm as Carbolic Acid—*B.M.J.* '97, i 972.

Stated to be best administered with Sugar of Milk, without a single bad result (*M.P.* '04, ii 523) in many hundred cases, including scarlet fever, diphtheria, croup, and cystitis.

Formic Aldehyde has received considerable attention as a therapeutic agent in pulmonary tuberculosis and has been employed both intravenously and as an inhalation. Maguire in his Harveian lectures recommends a solution of 1 part Formic Aldehyde Gas in 2000 parts of a sterilised solution of Sodium Chloride—*Trans. of Brit. Cong. on Tuberculosis*, vol. iii p. 438, *L.* '00, ii 1549, 1633, 1709, '01, i 629, 707, '01, ii 310, '03, i 98, '03, ii 463, *B.M.J.* '00, ii 1566, 1637, 1695.

Intravascular injections have shown that at present there is no evidence which would warrant the use of a septicæmia in animals can be favourably influenced by the intravenous injection of an antiseptic—*L.* '03, i 98.

A single intravenous injection of Formaldehyde in physiological solution effected a cure (*L.* '05, i 1341) in a case of marked oral sepsis, and the cure by intravenous injection in a case of tuberculous abscess of the lung in a patient with acute pulmonary tuberculosis was absolute and rapid. Tubercle bacilli, sputum and cough all disappeared.

Poisons such as Strychnine, Veratrine, Morphine, Atropine and Phosphorus will remain in the organs preserved with a 10 p.c. aqueous solution of Formalin for a very long time—*L.* '05, i 1093.

Formaldehyde has been somewhat extensively used by intravenous injection in the treatment of pulmonary phthisis, and a solution of 1 in 2000 is a suitable strength for injection.

As an inhalation, it has been used with good results in tuberculosis, pertussis and diphtheria. 2½ to 6 p.c. solutions of Formalin in pure Water or in 10 to 20 p.c. Glycerin solution are convenient, and should there be more than usual sensitiveness in the air passages, a little Aromatic Spirit of Ammonia may be added.

A solution of Formalin 1, Chloroform 1, Alcohol (90 p.c.) 2, has also been used, or it may be used as a fine spray at a strength of 6 to 10 p.c. solution mixed with Glycerin—*B M J* '99, 1 202, 772, 1440, '00, 1 189, '00, 1 1624, '02, 1 1692, *L* '01, 1 468, '01, 1 310, '02, 1 562, 772, *Trans of the Brit Cong on Tuberculosis*, vol III p 486.

Severe inflammation of the ends of all the finger nails caused by the prolonged use of a 1 in 500 solution of Formalin as a disinfectant for the hands—*B M J* '02, 1 54.

Recurrent papillomata of the larynx treated locally by Formalin as a 1 in 1000 increasing up to 1 in 100 spray—*L* '01, 1 487.

A solution of equal parts of Formalin and Glycerin as a paint in lupus—*B M J* '01, 1 1078, *B M J E* '01, 1 48.

10 to 50 p.c. ointment in chilblains if skin be not delicate—*P* '08, 1 251.

Cases of poisoning from swallowing commercial Formalin—*B M J E* '01, 1 9, 72.

A few drops of Liquor Ammonia Fort well diluted with Water, or still better Liquor Ammonii Acetatis, given at frequent intervals as an antidote in cases of Formalin poisoning—*B M J E* '01, 1 7.

Formic Aldehyde as a preservative of foods—It is generally condemned as a preservative of foods on account of its action on the flesh forming constituents, rendering them insoluble. The proteids of Milk containing Formalin fail to yield to the digestive action of Pepsin—*L* 99, 1 1507, '99, 1 1282, 1427, 1577, '00, 1 228, *J C S Abs* 01, 1 517, *B M J E* '02, 1 16.

Recommendation of the Departmental Committee appointed to inquire into the use of preservatives in food, that Formaldehyde or any of its preparations be absolutely prohibited in food or drinks—*L* '01, 1 1683, *B M J* '01, 1 1758, *P J* '01, 1 620, *C D* '01, 1 880, *Analyst*, '01, 383.

Formic Aldehyde as a disinfectant—There is no conflict of evidence as to Formaldehyde being a reliable disinfectant when used in solution, or used in the gaseous state for room disinfection when all objects are freely exposed, but it seems to be the general opinion that for the disinfection of heavy materials and furniture, or where there are many cracks or fissures, or the surfaces are not freely exposed, on account of its non-penetrative properties it is not so suitable as Sulphurous Acid Gas. It has the advantage, however, of being non-injurious to delicate fabrics such as furs, silks, etc.—*L* '99, 1 1436, '02, 1 759, '03, 1 37, *B M J* '99, 1 1280, '00, 1 1575, '00, 1 1600, '02, 1 792, *B M J E* '00, 1 55, *T G* '99, 600.

Report of the practical experiments on disinfectants undertaken by the London County Council both Formic Aldehyde and Sulphur Dioxide failed in the case of wood and cloth charged with spores, in the case of tuberculous sputum dried on linen and paper, Formaldehyde showed to greater advantage than Sulphur Dioxide—*L* '02, 1 759, *B M J* '02, 1 792.

Formaldehyde in the state of vapour is able to destroy the bacilli in 'dried' sputum, but solutions of 4 to 10 p.c. did not affect 'ordinary' sputum—*L* '03, 1 37.

A paper by Kanthack on the use of Formalin lamps for the disinfection of rooms—*L* '98, 1 1049.

The Aldehyde vapours are non-poisonous, but very irritating to the eyes and throat, they possess marked deodorant and disinfectant properties, and are well suited to the purposes of room disinfection, for they do not affect colours. The use of the reagent in a gaseous form appears to possess the advantages over disinfection by Sulphurous Acid, that it injures nothing except Iron, it diffuses better, and it possesses greater disinfectant power—*B M J* '98, 1 1542.

Muller's Fluid, containing 10 p.c. of Formal, has been recommended for hardening pathological specimens, but it deposits in 5 days and must be changed, 60 p.c. Alcohol, to which 1 p.c. Formal has been added, is a good preservative fluid after hardening in above—*B M J E* '96, 1 88.

The 35 p c solution is diluted with 10 to 50 of Water, for fixing and hardening histological and pathological specimens, and for preserving them.

In room disinfection, best results obtained when 50 grammes of Potassium Permanganate are added to 100 c c Formaldehyde, or multiples of these quantities. The space to be disinfected—*T G '07, 460*
value and selection of disinfectants—*P G '07, 269*

Foreign Trials. Official in Austr., Belg., Dan., Dutch, Fl., Ger., Ital., Jap., and U.S.

Tests.—Formic Aldehyde Solution has a specific gravity of 1.079 to 1.081. The *U S P* has a specific gravity of 1.075 to 1.081 at 25° C (77° F), the *P G* solution 1.079 to 1.081. It should be neutral or only faintly acid to Litmus paper.

On evaporating 5 c c to dryness on a water-bath a white amorphous mass is left, which should leave no weighable residue on ignition.

If the solution be made strongly alkaline with Ammonia Solution and evaporated to dryness on a water-bath, a white crystalline residue readily soluble in Water remains. Formaldehyde Solution readily reduces Silver Ammonio-nitrate Solution, and Potassio cupric Tartrate Solution, the former yielding a greyish-black deposit of metallic Silver, the latter a deposit of red Cuprous Oxide. 2 c c of the Solution mixed with an equal volume of Potassium Hydroxide Solution and about 0.5 gramme of Resorcin gradually yields, when the mixture is heated to boiling, a bright red coloration. 2 drops added to 5 c c of Sulphuric Acid containing a little dissolved Salicylic Acid yields, on permanent amount of brilliant blue colour, varying in amount of present, is produced when about 0.05 gramme of Phenylhydrazine Hydrochloride is added to 1 c c of the solution diluted to 5 c c with Distilled Water, followed by the addition of 8 drops of a freshly-prepared 5 p c Sodium Nitro-prusside Solution, thorough agitation of the liquid, then Sodium Hydroxide Solution drop by drop until an excess has been added. Numerous methods have been proposed for the quantitative determination of Formic Aldehyde. That perhaps most generally used, on account of the ease of manipulation is the Ammonia process, and depends upon its conversion into Hexamethylenetetramine and determination of the amount of Ammonia absorbed. A weighed quantity of 2 grammes of pure neutral Ammonium Chloride is dissolved in 25 c c of Water and introduced into a flask provided with a well-fitting stopper. A weighed quantity of 2.5 grammes of the sample is carefully neutralised with Normal Volumetric Potassium Hydroxide Solution and added to the Ammonium Chloride Solution. A measured quantity of 25 c c of Normal Volumetric Potassium Hydroxide Solution is then added, the flask securely stoppered and set aside for one hour. A few drops of Rosolic Acid Solution are added, and the excess of Ammonia is titrated with Normal Volumetric Sulphuric Acid Solution, each c c of Normal Volumetric Potassium Hydroxide Solution absorbed corresponding to 2 p c w/w of Formaldehyde.

The *P G* process consists in treating 5 c c of the Formic Aldehyde Solution with 20 c c of Water and 10 c c of Ammonia Solution, allowing the mixture to react for one hour in a well-stoppered flask. A measured quantity of 20 c c of Normal Volumetric Hydrochloric Acid Solution is added, a few drops of Rosolic Acid Solution and the excess of acid titrated with Normal Volumetric Potassium Hydroxide Solution. At least 4 c c should be necessary to produce a red coloration. The *U S P* process depends upon the oxidation of the Formic Aldehyde to Formic Acid and titration with standard alkali. A measured quantity of 3 c c of Formic Aldehyde Solution is placed in a well-stoppered flask and accurately weighed. A measured quantity of 50 c c of Normal Volumetric Sodium Hydroxide Solution is added, followed immediately by 50 c c of Hydrogen Dioxide Solution added slowly through a small funnel, a drop or two of Litmus Solution having been previously added, and the solution previously neutralised with Normal Volumetric Sodium Hydroxide Solution. When the reaction is completed the funnel and sides of the vessel are rinsed with Water. The whole allowed to stand 30 minutes, and titrated back with Normal Volumetric Sulphuric Acid Solution, using Litmus solution as an indicator. The number of c c of Normal Volumetric Sulphuric Acid consumed are subtracted from 50, the remainder is multiplied by 2.979 and the product divided by the weight of

Solution taken, the quotient indicates the percentage w/w of absolute Formic Aldehyde. Both methods have been tried in the author's laboratory. The Hydrogen Peroxide method is the more accurate of the two and yields higher results, but the ease of manipulation of the Ammonia process and the fact that the results yielded are sufficiently accurate for most practical purposes ensures its more general application. An Iodometric method has been proposed by G. Romijn (*Analyst* '97, 221). A weighed quantity of 2.075 grammes of Formaldehyde Solution is diluted with Water to 500 c.c. A measured quantity of 10 c.c. of this Solution is mixed with 25 c.c. of Deci-normal Volumetric Iodine Solution and sufficient Sodium Hydroxide Solution (15 p.c.) added drop by drop to colour the liquid clear yellow. Allow to stand 10 minutes and add sufficient dilute Hydrochloric Acid to liberate the uncombined Iodine which is titrated with Deci-normal Volumetric Sodium Thiosulphate Solution. 1 c.c. of the Volumetric Iodine Solution is equivalent to 0.001489 gramme of absolute Formic Aldehyde. A method based on the production of a Bisulphite compound by interaction between Formaldehyde, Sodium Bisulphite and Normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator, has also been suggested. The end reaction is, however, somewhat indefinite, and it is therefore difficult to judge when the reaction is complete. The method adopted by the *Fr. Codes* is similar to that of the *U.S.P.*, and depends upon the oxidation of the Formaldehyde to Formic Acid, by means of Hydrogen Peroxide, and determination of the Formic Acid volumetrically.

The more generally occurring impurities are Methyl Alcohol, excess of acid, e.g., Formic Acid, fixed impurities, Iron, Lead, Copper, and Calcium, Chlorides and Sulphates. The presence of Methyl Alcohol or Acetone may be shown by the Iodoform test, no precipitate of Iodoform should be produced when 1 c.c. of the Solution is mixed with 10 c.c. of Iodine Solution, the excess of Iodine decolourised with Sodium or Potassium Hydroxide Solution, and the mixture warmed, excess of acid may be determined by titration with Normal Volumetric Sodium or Potassium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, the *U.S.P.* and the *P.G.* permit 0.23 w/v of anhydrous Formic Acid, fixed impurities are detected by the residue left on evaporation and ignition, Iron by the Potassium Ferrocyanide test in the diluted solution, Lead, Copper, Calcium, Chlorides and Sulphates by diluting the solution 1 to 3 or 1 to 4 and applying Hydrogen Sulphide Solution, Ammonium Oxalate Solution, Silver Nitrate Solution or Barium Chloride or Nitrate Solution respectively.

PARAFORMIC ALDEHYDE (Paraform Tri-oxymethylene)—A white micro-crystalline or amorphous powder, insoluble in Water. It is a polymer of Formic Aldehyde, it volatilises at 100° C (212° F), and is readily convertible into that substance when heated to the above temperature in the presence of Water. It is used for disinfecting rooms. It is official in *Fr. Codes*.

Sterilisol—An aqueous solution of Paraform, the solution being effected at 40° to 45° C (104° to 113° F) *in vacuo*—*L.* '05, 1 1075

HEXAMETHYLENETETRAMINE—Colourless and odourless lustrous crystals, or as a white crystalline powder, possessing an alkaline reaction.

It is a condensation product, obtained by the action of Ammonia gas on Formic Aldehyde. It should be preserved in well stoppered bottles.

Commercial varieties of this substance are known under the names of **Aminoform**, **Cystamine**, **Formin** and **Urotropine**.

It has been implied, if not actually stated, that all these products are exactly the same and practically interchangeable. This is not borne out by clinical experience, for the different preparations do not always produce the same results in the same patient. This is precisely what we should expect from a body of which the stereographic formula presents so many different possibilities.

Solubility—Soluble 5 in 6 of Water, 1 in 8 of Alcohol (90 p.c.), sparingly in Ether.

Urinary antiseptic, given in cystitis and phosphaturia.

Dose.—5 to 15 grains = 0.32 to 1 gramme, dissolved in Water or in aerated Water.

Marvellous effects in doses of 10 grains thrice daily in typhoid bacilluria and cystitis, for which conditions it appears to be an almost specific remedy—*L* '00, 1 707, 1059, 1876, '01, 1 174, '02, 1 687, *B M J E* '02, 1 95 5-grain doses three times a day in cystitis with ammoniacal urine—*L* '00, 1 1658

As an intestinal disinfectant — *B M J E* '01, 11 60

In daily doses of 20 to 60 grains in diabetic coma.—*B M J E* '02, 1 72

Two cases of hæmaturia following the use of from 5 to 10 grains of the salt three times daily—*B M J* '01, 1473, 1617, 1659, *T G* '01, 617

In the pyuria of *tuberculosis* 3 grains daily — *L* '03, 11 1019

In vesical catarrh accompanying typhoid fever 10-grain doses have been given three times daily with advantage—*B. M. J.* '04, n. 70, p. 1705 (S). It is liable to produce irritation of the stomach, a measles-like rash, renal irritation, albuminuria and hematuria unless well diluted, and the powder of Water

In enteric fever, doubtful if it had any influence on the course of the illness itself — B M J 05, 1 414

In acute but not in tubercular cystitis, in 3-gramme doses — *T G '07*, 311

A mixture of Urotropine and Iridin has a pronounced effect in causing dissolution of calculi in the treatment of artificially-produced cholelithiasis.—*B M J* '05, ii 272

Of value as a prophylactic against the nephritis of scarlatina all cases of scarlatina be treated from the beginning with 5 to 10 grains daily, well diluted with Water, to be continued to the 28th day of the disease—*Edin Med Jour* '07, 1: 113

Foreign Pharmacopoeias - Official in Dan, Jap, Swiss and U S

Tests — **Phenol** : Dissolves in water at a temperature of 263° C (505.4° F). With concentrated Sulphuric Acid and heated with Sulphuric Acid, it develops a carmine-red coloration. If heated with diluted Sulphuric Acid the characteristic irritating odour of Formaldehyde is evolved, and if a piece of filter paper, moistened with Silver Ammonio-nitrate Solution, be held over the tube, it is immediately darkened. If this Sulphuric Acid Solution be cooled and supersaturated with Sodium Hydroxide Solution, the characteristic odour of Ammonia is evolved, recognised also by its turning a piece of moistened red Litmus paper blue. A 10 pc aqueous solution affords a precipitate with Tannic Acid Solution, with Mercuric Chloride Solution, on standing, white precipitates are produced, with Iodo-potassium Iodide Solution it yields a white precipitate.

The more generally occurring impurities are mineral matter, Copper, Iron or Lead, Chlorides or Sulphates, Ammonia salts, and Para formaldehyde. Mineral matter may be detected by the residue left on ignition. Copper and Lead may be detected by Hydrogen Sulphide Solution, Iron by Potassium Tartrate Solution, Chlorides by Silver Nitrate Solution after acidification with dilute Nitric Acid, Sulphates by Barium Chloride Solution after acidification with diluted Hydrochloric Acid. Ammonium salts and Parafomaldehyde may be detected by Potassium-mercuric Iodide (Nessler's) Solution, the former causing brownish-red colour or precipitate, the latter readily causing a separation of metallic Mercury.

AMYLOFORM—A white, amorphous, odourless powder, which is a compound of Formaldehyde with Starch Insoluble in Water, but when brought in contact with moist surfaces it is slowly decomposed, giving off Formaldehyde. Recommended as a dressing or as a dusting powder—*L* '97, 11 40, '00, 1, 470; *T G* '00, 816

Dextroform is a white powder, freely soluble in Water, slightly soluble in cold Glycerin, but dissolves in 10 when warmed. It is a compound of Formaldehyde with Dextrine. It has been used internally, and has been given in the form of a 5, 10, or even 20 per cent solution in gonorrhoea.

Glutol is a yellowish-white powder, insoluble in Water and Glycerin; it is a compound of Formaldehyde with Gelatin, used as an antiseptic dressing.

FORMICIN—A syrupy liquid, sp gr 1.240 to 1.260. Miscible with Water, Alcohol and Chloroform in all proportions. It is produced by the action of Acetamide on Formaldehyde, and has been introduced (*B M J E* '05, ii. 99, *P J* '05, ii. 885) as a powerful antiseptic. Applied in the form of a 2 p.c. tepid solution it has been used as a surgical disinfectant.

HELMITOL (Hexamethylenetetramine Anhydromethylene Citrate)—Colourless crystals, or as a white crystalline powder, soluble 1 in 5 of Water, sparingly soluble in Alcohol (90 p.c.), insoluble in Ether. Has been recommended in chronic posterior urethritis, cystitis and prostatitis.

In acute cystitis the subjective results were good (*B M J E* '05, ii. 20), but the cystitis reappeared if the drug were discontinued. It is more useful in bacteriuria, the results being permanent.

Dose—10 to 15 grains = 0.65 to 1 gramme, three times daily.

HETRALIN (Dioxybenzolhexamethylenetetramine)—One of the many derivatives of Hexamethylenetetramine, it forms snow-white crystals, soluble (according to our experiments, *P J* [4], xx. 784, *C D* '05, i. 788) 1 in 9 of Water, 1 in 17½ of Alcohol (90 p.c.), 1 in 180 of Ether, sp gr 0.785, insoluble in Chloroform. It has been introduced as a urinary antiseptic (*B M J* '04, ii. 1468), but (*B M J E* '04, ii. 64) failed to produce any good effect in seven cases of tuberculous disease of kidneys and bladder.

In doses of 1 grain, given every 3 hours, has been found of value in acute cystitis in an infant—*B M J* '07, i. 1181.

In cases of cystitis it has proved beneficial in doses of 15 grains twice a day in a tumblerful of cold Water—*General Practitioner*, Feb. 18, 1907.

INDOFORM—A white powder, m.p. 108° to 109° C (226.4° to 228.2° F.), produced by action of Formaldehyde on Acetyl-salicylic Acid. Sparingly soluble in cold Water, and has an acid, astringent taste. Antirheumatic and anti-neuralgic.

Dose—7½ grains = 0.5 gramme.

SODIUM ANHYDROMETHYLENECITRATE (Citarin)—A white granular, amorphous powder, soluble 1 in 1½ Water, insoluble in Alcohol (90 p.c.), and in Ether. Given in rheumatism and gout, and as a solvent for Uric Acid calculi.

Dose—15 to 30 grains = 1 to 2 grammes.

URESIN (Hexamethylenetetramine Di-lithium Citrate)—A white, crystalline powder, readily soluble in Water, has been given in gout, and as a solvent for certain urinary deposits.

Dose—5 grains = 0.32 gramme.

Chinotropine (Quinotropine) is a white powder, readily soluble in Water. It is a combination of Quinic Acid and Hexamethylenetetramine. Is said to lessen formation of Uric Acid—*B M J E* '01, ii. 95, *P J* '01, i. 666.

Dose—10 to 15 grains = 0.65 to 1 gramme.

Under the name of Igazol a combination of Formic Aldehyde with Chloral, Terpene and Iodoform has been introduced for the treatment of pulmonary consumption, and is used as an inhalation—*Trans of Brit Cong on Tuberculosis*, iii. 416, *B M J* '00, ii. 662.

Lysoform is a clear, colourless or pale yellowish, soapy liquid. Miscible with Water. Introduced as an antiseptic. A solution 1 to 2 tablespoonfuls to the pint is used to disinfect the hands—*B M J E* '01, ii. 88, *L* '03, ii. 1807.

In the sterilisation of the hands, a 2 p.c. solution in Alcohol gave much superior results to the hot Water-Alcohol method (*B M J* '05, i. 727), but still better results were obtained with Bacillol and Sublimin. Bacillol is a Cresol preparation, non-toxic and non-injurious to the hands. Experiments with a 1 p.c. alcoholic solution gave complete sterility through all tests in a minimum of 60 p.c.

Carbol Lysoform is stated to be a mixture of crude Carbohc Acid and Lysoform, and to be a more active bactericide than either of its components—*B M J E* '02, ii. 92, *P J* '03, i. 840.

Not Official.

FUCUS VESICULOSUS

Bladder-wrack collected from rocks by the seaside and dried

Medicinal Properties—Given to reduce obesity
Smelling fresh seaweed is said to relieve hay asthma

Foreign Pharmacopœias—Official in Mex (Encina de Mai), Port (Bodelha) Not in the others

Descriptive Notes—This seaweed is of a blackish colour, flat, forked, about half an inch broad and a foot or more long. It is distinguished by having a mid-rib, and oval air bladders in the frond, usually in pairs, one on each side of the mid-rib. When dried it has the essence of Mannite on the surface. It is said to be used in pills.

Tests—Bladder-wrack leaves about 15 p.c. of ash. Two specimens examined in the author's process of air. The ash is blackish and contains 15.6 p.c. of ash.

EXTRACTUM FUCI VESICULOSI—Prepared from the dried seaweed. Alcohol (45 p.c.), and evaporation to a stiff extract, incorporated in the B.P.C. under the title *Extractum Fuci Vesiculosi*. Since 1867, and was used in pills.

Dose—3 to 10 grains = 0.2 to 0.65 gramme.

Test—It leaves about 18 p.c. of ash on ignition. The ash has a specific gravity of 1.044.

EXTRACTUM FUCI VESICULOSI LIQUIDUM—Extract of Fucus Vesiculosus in Alcohol (45 p.c.) to saturation, incorporated in the B.P.C. under the title *Extractum Fuci Vesiculosi Liquidum*. The fluid extract has been given in *Companion* and *Formulary* '01.

Dose—1 to 2 fl. drms = 3.6 to 7.1 c.c.

Tests—Liquid Extract of Fucus Vesiculosus in Absolute Alcohol (45 p.c.) to saturation, contains about 8 p.c. w/v of solid matter. The solid matter is held in suspension by a piece of gelatin.

GALBANUM

GALBANUM

FR, GALBANUM, GER, GALBANUM, ITAL, G

A Gum-resin obtained from *Ferula galbanifolia* and probably from other species.

Gambanum contains about 9.5 p.c. of impurities and Buhse, Gum-resin soluble in Alcohol, and 27 p.c. of impurities. The pure Gum-resin contains Umbelliferone Galbanol, about 50 p.c., and similar Ester, 20 p.c., Umbelliferone, 0.5 to 3.0 p.c. of ash.

Medicinal Properties—Internally, less energetic, externally as a plaster, in chronic inflammatory swellings.

Dose—5 to 15 grains = 0.32 to 1.0 gramme

Official Preparation—Pilula Galbani Composita

Not Official—Emplastrum Galbani and Unguentum Galbani Compositum

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed and Swiss. Not in Hung or U.S.

Descriptive Notes—Galbanum is much scarcer in commerce than formerly. Two principal varieties of the drug are recognised in commerce, which are called respectively Levant and Persian, although both are the products of Persia. The Levant Galbanum, which comes by way of Egypt and Turkey, occurs in two forms (1) small yellowish-brown tears, yellowish-white and opaque internally, and possessing a musky odour and bitter and somewhat acrid taste, and probably obtained from the stem, (2) a tough, pasty mass, consisting of slices of root with bluish green, almost translucent, pieces, mixed with yellowish-brown pieces, and also possessing a musky odour, and evidently obtained from the root. Both of these probably come from near Shiraz, *via* the Persian Gulf. The Persian Galbanum occurs also in two forms (1) a turpentine, sticky mass, having a turpentine rather than a musky odour, and containing fruit stalks, but no slices of roots, (2) a treacly liquid, of a reddish colour, often containing fruits of the plant. These apparently come from the Demawend mountains in the north of Persia, by way of Astiakhian and Orenburg, and are apparently the produce not of *F. galbaniflua* but another species. African Ammoniacum, the only Gum-resin that at all resembles Galbanum, does not yield Umbelliferone. Persian Galbanum gives a yellowish-red colour with Hydrochloric Acid, whilst the Levant gives different shades of violet. As the former possesses a musky odour, and the latter a turpentine one, they are probably derived from different species. The *P. G.* directs Galbanum to be dried over quicklime and submitted to a low temperature in order to powder it.

Tests—Galbanum yields about 50 p.c. of substances soluble in Alcohol (90 p.c.). If a portion is heated to redness in a dry test-tube, the residue, when cooled and boiled with Water, yields a solution which, largely diluted, produces a strong blue fluorescence when rendered alkaline with Ammonia Solution. This test is known as the Umbelliferone test, and remarks upon its application will be found under Ammoniacum. The ash should not exceed 10 p.c. The volatile Acid value is 73.5 to 114.0, the Acid value, 21.2 to 63.5, the total Saponification value, 116.2 to 135.8.

Ammonia.—If finely powdered Galbanum be boiled with fuming Hydrochloric Acid for a quarter of an hour, filtered through a previously moistened filter, and the filtrate carefully saturated with Ammonia Solution, the mixture shows a blue fluorescence in reflected light.—*P. G.*

Residue from Alcohol (90 p.c.).—After completely exhausting 100 parts of Galbanum with boiling Alcohol (90 p.c.), a residue is obtained which, after drying, should amount to at most 50 p.c. of the original mass.—*P. G.*

Ash.—100 parts of Galbanum should yield on incineration not more than 10 parts of ash.—*P. G.*

Preparation

PILULA GALBANI COMPOSITA — COMPOUND PILL OF GALBANUM *B. P. Syn* — COMPOUND PILL OF ASAFETIDA

Asafetida, 1, Galbanum, 1, Myrrh, 1, Syrup of Glucose, *as* Mix together on a water-bath

Dose—4 to 8 grains = 0.26 to 0.52 gramme

The following modification will be found convenient for dispensing powder the Myrrh, mix it with the Asafetida and Galbanum melted on a water-bath, allow the mixture to cool, and after crush it by attrition in a mortar to powder with one-sixth of its weight of Light Magnesium Oxide. The powder will keep well, and can be made into pills as required with the aid of Alcohol (60 p.c.)

Foreign Pharmacopœias—Official in Port, similar to Brit Not in the others

Not Official

EMPLASTRUM GALBANI—Galbanum, 1, Ammoniacum, 1, melt together and strain, then add them to Yellow Beeswax 1, Lead Plaster 8, previously melted together Mix (1 in 11)

Was official in L.P. 1835, but omitted in B.P. 1898, it has been incorporated in the B.P.C. using 7 of Lead Plaster instead of 8 so as to make the total 10

A plaster more or less resembling this is Official in the Foreign Pharmacopœias except Hung and U.S. Fr has 1 plaster of dissection for the Mex Emplastro de Galbano Azafranado Dutch Emplastrum Galbani Compositum

UNGUENTUM GALBANI COMPOSITUM—Galbanum Plaster, 4 oz, Lead Plaster, 4 oz, White Beeswax, 4 oz, soft Extract of Opium, 1 dram, Olive Oil, 20 fl oz Melt together

It is used for boils and carbuncles, and for sore nipples and inflamed breasts

GALLA.

GALLS

FR, GALLE D'ALEP, GER, GALLAPPEL, ITAL, NOCI DI GALLA, SPAN, AGALLA DE ALEPO

Excrecences on *Quercus* Oliv, resulting from punctures and deposited eggs of *Galleria mellonella*, Oliv

This description occurs in B.P. and U.S.P. Fr Codex gives it as a pathological product due to the puncture of *Cynips* Oliv, or the young shoots of the oak of the dyers, *Quercus tinctoria*, Lamk. var *infectoria*, Oliv

Chiefly from Turkey, Persia and Greece

Galls contain 60 to 70 p.c of Gallo-tannic Acid, and 3 to 5 p.c of Gallic Acid, to which their therapeutic qualities may be attributed

Solubility—All the soluble matter of Galls is taken up by forty times their weight of boiling Water, and the residue is tasteless

Medicinal Properties—Astringent Chiefly used locally in form of lotion or injection to suppress hæmorrhage from the gums, nose, etc., to lessen the discharge from mucous membranes, as in gleet, leucorrhœa etc., both Ointments are useful in painful hæmorrhoids

Dose—10 to 20 grains = 0.65 to 1.3 gramme

Incompatibles—The mineral Acids, Iron and Lead salts, Copper Sulphate, Silver Nitrate, Potassium and Sodium Carbonates and Alkalies, Lime Water, Tartar Emetic, Ipecacuanha and Opium, Infusions of Cinchona, Calumba and Cuscuta

Official Preparations—Unguentum Galla and Unguentum Galla cum Opio Used in the preparation of Acidum Gallicum and Acidum Picricum

Not Official—Decoctum Galla, Suppositoria Galla, and Suppositoria Galla cum Opio

Foreign Pharmacopœias—Official in Austr, Dan, Dutch, Fr (Galle d'Alep), Ger, Hung, Ital (Noci di Galla), Jap, Mex (Agallas de Levante), Norw, Port (Galha), Russ, Span (Agalla de Alepo), Swiss and U S

Descriptive Notes—The galls of *Quercus infectoria* are known in commerce as Aleppo galls, and are met with in three varieties, blue, green, and white. The blue green are considered the best, the dark green second, and the white galls are of very inferior quality. The last named, besides the pale yellowish-brown colour, are noticeable for the fact that each shows a perforation whence the gall insect has escaped. They are also lighter in weight, and are excluded from use by the official description, according to which Aleppo galls are spherical, averaging $\frac{1}{2}$ to $\frac{3}{4}$ in (12 to 18 mm) in diameter, and have a smooth surface, are dark green or olive green externally, are furnished in the upper half with small pointed tubercles, and ridges widely separated, the lower half, being usually smooth, are yellowish or brownish white internally, with a small central cavity. They have an astringent and slightly acid taste, followed by a slight sweetness. The characteristic features of powdered galls are the raphides, angular fragments of Tannin, the parenchymatous cells, with intercellular spaces, the sclerenchymatous cells with stratified walls, and starch grains with a stellate hilum. English oak galls, from *Quercus pedunculata*, Willd, resemble Aleppo galls in size, but have no prominences, and contain less than a third of the amount of Gallic tannic acid (15 to 20 p c) contained in the Aleppo galls (70 p c). Other oak galls, under the name of Morea galls, are occasionally imported from Greece. These are about $\frac{1}{2}$ in in diameter, and have a crown of small tubercles. The Japanese and Chinese galls, from Hiogo and Canton, which are largely imported, are irregularly fig shaped, hollow, and downy externally, from 1 to 2 in long, $\frac{1}{2}$ to 1 in broad, the shell being only $\frac{1}{16}$ to $\frac{1}{12}$ in in thickness. They are formed on *Rhus semialata*, Murr, and other species by *Aphis chinensis*, Bell, the skeletons of which are usually found within the galls. They yield up to 78 p c of Gallic tannic Acid, and are therefore of considerable technical value. The plum-shaped Chinese galls are formed on *Distylium racemosum*, S et Z, Tamarisk galls, formed on *Tamarix orientalis*, L, and other species, are from the size of a pea up to $\frac{1}{2}$ in in diameter, and are occasionally imported, they contain about 40 p c of Tannin.

ACIDUM GALLICUM—See ACIDUM GALLICUM

ACIDUM TANNICUM—See ACIDUM TANNICUM

Preparations

UNGUENTUM GALLÆ—GALL OINTMENT

Galls, 1, Benzoated Lard, 4 (1 in 5)

Foreign Pharmacopœias—Official in U S, 1 in 5 Not in the others

UNGUENTUM GALLÆ CUM OPIO. GALL AND OPIUM OINTMENT

Opium, $7\frac{1}{2}$ grains, Gall Ointment, 92 $\frac{1}{2}$ grains (about 1 in 13)

The ointment might be made direct by mixing 15 grains of Opium and 37 grains of Galls with 118 grains of Benzoated Lard

Not Official

DECOCTUM GALLÆ.—Bruised Galls, 2½, Distilled Water, 40; boil to 20, and strain (1 in 8)

B.P.C. Decoction is 1 in 16

SUPPOSITORIA GALLÆ.—5 grains powdered Galls and 1 grain Opium in each, with a basis of Coconut Stearin

TINCTURA GALLÆ.—1 of Galls percolated with Alcohol (60 p.c.) to yield 8, (1 in 8)

Dose —½ to 2 fl drms = 18 to 71 c.c.

B.P.C. Tincture is 1 in 10, also with " " " " " "

Foreign Pharmacopœias—Official in Austr., Dan., Dutch, Ger., Hung., Jap., Mex., Norw., Russ., Swiss and U.S., 1 in 5. All by weight, except U.S. Not in the others.

Not Official.

GARCINIA PURPUREA, Roxb.

KOKUM BUTTER TREE

Grows in the forests of Malabar, the Concars, and other parts of the Madras Peninsula

The Oil of the seeds (*Kokum Butter*) is obtained by first exposing the seeds for some days to the action of the sun to dry, they are then bruised and boiled in Water, the Oil collects on the surface, and on cooling contracts into a solid cake. It melts at 98° F. (36.6° C.). The seeds yield about 10 p.c. of Oil.

It is used in India in the preparation of ointments, suppositories, etc.

Not Official

GAULTHERIÆ OLEUM.

OIL OF WINTERGREEN.

Three nearly allied substances are sold as Oil of Wintergreen, and they are all official in U.S.

Oil of Gaultheria (Wintergreen)—A volatile Oil distilled from the leaves of *Gaultheria procumbens*, L., consisting almost entirely of Methyl Salicylate, and nearly identical with Volatile Oil of Betula. A colourless or yellow liquid, with a strong characteristic odour, and a pungent taste.

Official in the *Ind.* and *Col. Add.* for the North American Colonies.

It should be kept in well-closed bottles of a dark amber tint, in a cool atmosphere, and protected as far as possible from the light.

It contains, according to Power and Kleber, about 99 p.c. of Methyl Salicylate, with a small amount of a paraffin, probably triacontane, an aldehyde or ketone, an apparently secondary Alcohol and an Ester.

Tests—Oil of Gaultheria has a sp. gr. of 1.175 to 1.185. It is slightly laevogyrate, the optical rotation being not below -0.25° nor more than -1° in a 100 mm tube. It boils at 218° to 221° C. (424.4° to 429.8° F.). It should form a perfectly clear solution at about 20° C. (68° F.) with 5 parts of Alcohol (70 p.c.). It should yield the tests and be free from the impurities mentioned under Methyl Salicylate. It may be distinguished from Oil of Betula by its optical rotation, the latter being optically inactive. Foreign oils of Gaultheria, if present, may be determined by the sp. gr.,

Volatile Oil of Betula (Sweet Birch) —A volatile Oil obtained by distillation from the bark of *Betula lenta*, L. It is identical with Methyl Salicylate, and nearly identical with Oil of Gaultheria

It should be kept in well closed bottles of a dark amber tint, in a cool atmosphere, and protected as far as possible from the light

It is produced by the action of the ferment Betulase on the glucoside Gaultherin

According to Power and Kleber, the Oil consists, to the extent of about 99.8 p.c., of Methyl Salicylate, and in its unrectified state of a paraffin, probably Triacotane and an Ester, but does not contain the secondary Alcohol found in Gaultheria Oil

Tests —Oil of Sweet Birch has a sp. gr. of 1.180 to 1.187. It is optically inactive. It boils between 218° and 221° C (424.4° and 429.8° F). It should form a perfectly clear solution at about 20° C (68° F) with 5 parts of Alcohol (70 p.c.)

Foreign Oils or Petroleum, if present, may be detected by a lowering of the sp. gr. In other respects it resembles Oleum Gaultheriae, and conforms to the tests and should be free from the impurities mentioned under that Oil

Methyl Salicylas ($\text{CH}_3\text{C}_6\text{H}_4\text{O}_2$, eq. 150.92) is produced synthetically. A colourless or slightly yellowish liquid, with a characteristic odour and taste. A large proportion of the Oil in commerce is synthetic Methyl Salicylate, or Artificial Oil of Wintergreen

It should be kept in well closed bottles of a dark amber tint, protected as far as possible from the light and in a cool atmosphere

Solubility —Readily soluble in Alcohol (90 p.c.), Ether, Chloroform, and Glacial Acetic Acid, only slightly soluble in Water

Medicinal Properties —A valuable remedy in acute rheumatism, internally, also externally, applied directly over joints and limbs and covered with oiled silk or gutta percha tissue, to prevent evaporation, thus applied is specially useful in acute muscular rheumatism, also mixed with equal parts of Olive Oil. Used largely as a flavouring agent in America, more particularly in dentifrices. It is a good antiseptic

Methyl Salicylate is better for external application than the Oil of Wintergreen as it does not produce an eruption. In all cases it was applied according to the process, become classic, of 50 to 100 drops poured upon a double fold of aseptic gauze, and covered by an impermeable material, applied for some hours, either to the forearm or to the leg, and renewed twice every twenty-four hours. The part treated with natural essence of Wintergreen, was more or less red, painful, and covered sometimes with a rubeoliform eruption, pure Methyl Salicylate produced no such reaction —*L* '98, i 52, *B M J E* '00, i 56

As a dressing in the treatment of chorea, 6 to 10 grammes of the Oil either pure or mixed with Vaseline and covered with oiled silk to prevent evaporation —*T G* '99, 240, *B M J E* '99, i 8

In subacute and chronic rheumatism it is stated to be of great advantage, employed either alone or in conjunction with Sodium Salicylate —*T G* '99, 612, *B M J E* '99, i 63

Dose —5 to 15 minims = 0.3 to 0.9 c.c. every 4 hours when given as a substitute for Sodium Salicylate, but the taste is rather pungent

Prescribing Notes —When required to be added into an emulsion or pills, the same general rules would apply as for other Essential Oils, see 'Mucilage Acaciae' and 'Pilulae'; or it may be given in Coffin's rules, containing 5 or 10 minims in each

Foreign Pharmacopœias —Official in Fr. and U.S. Not in the others

Tests —Methyl Salicylate has a sp. gr. of 1.185 to 1.190. It is optically inactive. It has a boiling point of 219° to 221° C (426.2° to 429.8° F). It is readily soluble in Alcohol (90 p.c.), the solution being neutral or only slightly acid to Litmus paper. *Fr. Codex* gives the sp. gr. as 1.1819 at 18° C (60.8° F), and the boiling point as 224° C (435.2° F)

The saturated aqueous solution yields with Ferric Chloride T.S. a deep violet coloration. It should form a perfectly clear solution at about 20° C (68° F) with 5 parts of Alcohol (50 p.c.). If the Oil be saponified with Sodium Hydroxide Solution and the alkaline liquid be filtered with about 10 c.c. of its volume of Water, and acidified with diluted Sulphuric Acid, a white crystalline precipitate is formed, which, collected on a filter, washed with a little Water, and recrystallised from hot Water, should possess a melting point of 150° to 157° C (312° to 314.6° F) and should otherwise answer the tests of identity of Methyl Salicylate from the impurities mentioned under Acidum Salicylicum. It may be determined volumetrically with Normal Volumetric Potassium Hydroxide Solution, using Phenolphthalein Solution as an indicator. A weighed quantity of 5 grammes of the Oil is dissolved in 25 c.c. of the Normal Volumetric Solution and the mixture is boiled for five minutes to effect saponification. It is cooled, and the excess of alkali is titrated with Normal Volumetric Sulphuric Acid Solution 1 c.c. of Normal Volumetric Potassium Hydroxide Solution corresponds to 0.15092 gramme of Methyl Salicylate. The number of c.c. of Normal Solution absorbed multiplied by 0.15092 and the product multiplied by 20, yields the percentage w/w of absolute Methyl Salicylate present in the sample. The percentage of Methyl Salicylate may also be determined by saponification with Normal Volumetric Potassium Hydroxide Solution or, adding sufficient Normal Volumetric Hydrochloric Acid Solution to produce a faintly acid reaction, removing the liberated Salicylic Acid by Ether, washing till free from mineral acid and titrating the ethereal solution of Salicylic Acid with Normal Volumetric Potassium Hydroxide Solution using Phenolphthalein Solution as an indicator. 1 c.c. of Normal Volumetric Potassium Hydroxide Solution corresponds to 0.15092 gramme of absolute Methyl Salicylate. A good specimen contains not less than 99 p.c. w/w of Methyl Salicylate.

The more generally occurring impurities are Alcohol or Chloroform, other volatile oils or Petroleum and Methyl Benzoate. Alcohol or Chloroform may be detected by placing 1 c.c. in a flask provided with a suitable condenser and heating on a water bath. The distillate should not have the characteristic odour of Alcohol or Chloroform. The presence of volatile oils or Petroleum is indicated by the separation of oily drops either on the surface or at the bottom of the liquid, when 1 c.c. of Methyl Salicylate contained in a capacious test-tube is agitated with 5 c.c. of Potassium Hydroxide Solution. Methyl Benzoate is indicated by the m.p. of the acid, obtained after saponification and the decoloration of the Salicylic Acid as described above.

SPIRITUS GAULTHERIÆ—Oil of Gaultheria, 5, Alcohol (95 p.c.), 95, both by measure—L.S.P.

Average Dose—30 minims. This has been incorporated in the B.P.C. using Alcohol B.P. in place of Alcohol L.S.P.

SANOFORM (Di-iodomethyl salicylate)—A white crystalline powder, almost odourless and tasteless. It contains 62.7 p.c. of Iodine.

The m.p. of the powder is 110.5° C (230.9° F), and it therefore may be sterilised at 100° C (212° F) without decomposition. It may be employed (B.M.J.E. '05, 1, 80) in all cases where Iodoform is used, chiefly as a dressing in minor surgical operations, in cases of senile and diabetic gangrene and in gynaecology.

Solubility—Insoluble in Water and Glycerin, slightly soluble in cold Alcohol (90 p.c.), and readily in Ether.

MESOTAN (Salicylic Acid Methoxymethylester)—A yellow, oily liquid, possessing a slight aromatic odour. Insoluble in Water, readily soluble in Alcohol (90 p.c.), Ether and Chloroform. It is stated to be readily absorbed by the skin, and to be useful as a local application in all forms of rheumatic and gouty affections. It may be used as a 50 p.c. solution in Olive Oil, or by itself—B.M.J.E. '03, 1, 44.

Applied externally as an antirheumatic, it is stated (L. '04, 1, 100) B.M.J.E. '05, 1, 20) to have afforded distinct relief. It is stated to afford relief in the after-treatment of acute rheumatism, but is not so during the fever. Not the slightest effect followed its use in chronic rheumatism, and in one case well-marked local effect.

There can be little doubt (*B M J* '05, i 715) of the value of Mesotan in the treatment of rheumatism, but its use requires caution and careful supervision. An embrocation consisting of equal parts of Mesotan and oil very gently applied to the feet and ankles caused a rash in about 10 or 12 days, not only on the parts to which the embrocation had been applied, but also on the arms. It should be painted on where the skin is specially delicate (*M P* '05, i 452), and the skin should be previously dried and not covered with any impervious material afterwards.

Methyl-acetyl-salicylate is a crystalline powder, insoluble in Water, soluble in Alcohol (90 p c) and in Chloroform. Has been recommended in rheumatic affections—*C D* '03, ii 90.

Amyl Salicylas—A colourless or slightly yellowish liquid, with a characteristic odour and taste. It is not nearly so pungent as Oil of Gaultheria, and therefore has been suggested as a substitute for the latter.

GELATINUM.

GELATIN

The purified air-dried product of the hydrolysis of certain animal tissues as skin, ligaments and bones by the action of boiling Water.

Commercial Gelatin varies considerably in its gelatinising power, and it is advisable to keep to the same brand to avoid alteration in formulas.

Medicinal Properties—Hæmostatic. Used for increasing the coagulability of the blood in aneurisms.

A sterilised 1 to 2 p c solution in normal saline has been used with considerable success in the treatment of aortic aneurism—*L* '02, ii 169, 558, '03, i 591, 1810, *B M J* '03, i 1493. *B M J E* '02, i 16, 91, *P* lxii 577, *P J* '99, ii 213, *C D* '01, ii 442.

A method of preparing the sterilised solution in flasks, and a description of a suitable apparatus for its use—*B M J* '01, 1415.

Each flask contains the requisite quantity of sterile Gelatin solution ready for use without further dilution and consequent risk of contamination. 100 c c of a 2 p c solution are introduced (*L* '05, i 1169) into the subcutaneous tissue over an interval of from ten to twelve minutes in order to avoid discomfort and over-distension of the skin. The inner aspect of the thigh is found to be the most convenient place for the injection. Potassium Iodide in 10 grain doses three times a day is given concurrently with the Gelatin injection.

Rectal injection of 250 c c of a 5 p c sterilised aqueous solution of Gelatin in the treatment of hæmoptysis—*L* '03, i 578.

Six samples of Gelatin examined, and tetanus spores found in four of them—*L* '03, i 579.

Cases of tetanus terminating fatally following the subcutaneous injection of Gelatin solution—*B M J* '01, ii 638, 741, *L*, '03, ii 33, *C D* '01, ii 382.

In melæna hæmatorum—*B M J* '02, i 43.

Contra-indicated in nephritis—*B M J E* '00, ii 1.

In hæmoptysis as a rectal injection, $\frac{1}{2}$ pint (*B M J* '05, i 68) three times daily.

Following frequent references to the use in aneurism of a 2 p c solution in normal saline injected into the gluteal region the further note appears. A popliteal aneurism was cured by seven injections, into the gluteal region, of a serum containing 2 p c of Gelatin. Each dose was 200 grammes, and the injections extended over seven months—*L* '05, i 1169.

Preferable to begin with 50 c c of 2 p c solution and gradually to increase the quantity to 100 c c and the strength to 4 p c. Treatment to continue over three to six months—*B M J E* '06, ii 76.

Official Preparations—Used in the preparation of the various Lamellæ, and Suppositoria Glycerini, p 569

Not Official—Gelatin Basis for Pessaries and Suppositories, Glyco-gelatin, Gelato-glycerin and Gelatinum Glycerinatum

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung., Jap, Mex, Port, Russ, Span, Swed, Swiss and U S Not in the others. Fr has Gelatine, and Gelatine officinale Swed includes a white and an ordinary Gelatin

Tests.—Gelatin, when immersed in cold Water, swells up and softens, taking up from 5 to 10 times its weight of Water, without undergoing solution to any great extent. It dissolves readily in hot Water. It is official in 1 in 50 parts of the latter liquid, a solution which is inodorous and which solidifies to a jelly on cooling. The *USP* states that a solution 1 in 50 of boiling Water should solidify on cooling and form a firm jelly; the *PG* states that a 1 in 100 solution will form a jelly on cooling. A useful test for comparing the gelauminising power of commercial Gelatins is to place 5 grains in a test-tube ($\frac{1}{2}$ in in diameter) with 250 grains of Water for half an hour, warm gently until the water is then place the test-tube in Water at 15°C (60°F), and leave undisturbed for 30 minutes, by which time a jelly should be formed of such consistence that it will remain in position if the test-tube be inverted. The aqueous solution affords a whitish precipitate with Tannic Acid Solution, *USP* specifies the strength of Solution as 1 in 5000, the *PG* says 'very dilute solutions'. Mercuric Chloride also affords a precipitate in an aqueous solution, neither Ferric Chloride TS, Lead Acetate Solution nor Alum Solution produces a precipitate. Potassium Bichromate Solution added to a Solution of Gelatin in hot Water, forms, on cooling, a jelly which becomes insoluble in warm Water after exposure to light. This latter reaction is made use of in photo-lithography. The *USP* and *PG* require that when incinerated it shall not leave more than 2 p.c. of ash.

Not Official

GELATIN BASIS FOR PESSARIES AND SUPPOSITORIES—Soften 1 oz. of Gelatin by allowing it to soak in 1 fl. oz. of Water until it is absorbed, then dissolve in $3\frac{1}{2}$ fl. oz. of Glycerin by the heat of a water-bath and allow it to cool and solidify. It can be medicated by mixing it over a water-bath and suspending or dissolving in it substances in fine powder, and then pouring the mixture into moulds.

See also Glycerin

GLYCO-GELATIN—Refined Gelatin 1 oz., Glycerin (by weight), $2\frac{1}{2}$ oz.; Ammoniacal Solution of Carmine a sufficiency, Orange-flower Water, $2\frac{1}{2}$ fl. oz.—Throat

Soak the Gelatin in the Water for 2 hours, then heat in a water-bath till dissolved, add the Glycerin and stir well together. Let the mixture cool, and when nearly cold add the Carmine solution, mix all uniformly coloured, and set aside to solidify.

This mass is used for making the various medicinal pills. The various substances are rubbed with an equal quantity of hyssop leaves when melted over a water-bath.

Glyco-gelatinum.—Gelatin, 12, Glycerin, 20, Sugar, 5, Citric Acid, 2, Carmine, a sufficient quantity.—*B P C*

This mass is stated to be of an unsatisfactory consistence, the following is an improvement (*P J '07*, n 804, 813) —

Gelatin, 20, Glycerin, 80, Distilled Water, 56, Orange-flower Water (undiluted), 7, Citric Acid, 2 50, Absolute Alcohol, 1, Oil of Lemon, 0 20, Solution of Carmine, 1

GELATO-GLYCERIN — Refined Gelatin (by weight), 5 oz, Glycerin (by weight), 6 oz, Water (by weight), 6 oz Soak the Gelatin in the Water for 12 hours, with occasional stirring, add the Glycerin, dissolve in a water-bath, and evaporate to produce 15 oz by weight of the Gelato glycerin — *Throat*.

(For preparing Nasal Bougies)

This has been incorporated in the *B P C*

GELATINUM GLYCERINATUM — Gelatin, 1, Glycerin, 1, Water, quantity sufficient to make 2 (all by weight) — *U S P*

Cover the Gelatin with boiled and Distilled Water, and after one hour drain the excess of Water away, transfer to a tared dish, add the Glycerin, and heat on a water-bath until solution is effected, strain whilst hot, and evaporate to 2

GELSEMII RADIX.

GELSEMIUM ROOT

FR, GELSEMIUM, GER, GELSEMIUMWURZEL, ITAL, GELSEMIO,
SPAN, GELSEMIO

The dried Rhizome and Roots of *Gelsemium nitidum*, Michaux

The plant, Carolina Jasmine, grows in the Southern States of North America

The root contains two alkaloids Gelsemine and Gelseminine It also contains B methylasculetin, which is identical with Gelsemic Acid

Excellent papers on the alkaloidal content of Gelsemium Root and Rhizome appear in *Proc Amer Pharm Assoc*, lvi 282, liv 383

Medicinal Properties.—Antispasmodic and analgesic Has been used in dental neuralgia, migraine, and especially in tic-douloureux (neuralgia of fifth nerve), also in uterine and ovarian pain, spasmodic and asthmatic cough, and in chorea

This drug should be used with care, and in the event of toxic symptoms presenting themselves, artificial respiration should be carried on — *Pr* 15 50

Official Preparation — Tinctura Gelsemii

Not Official — Extractum Gelsemi Alcoholicum, Fluidextractum Gelsemi

Antidotes — Emetic of Mustard and Water, Atropine, Aromatic Spirit of Ammonia, Brandy, Nitroglycerin, and Digitalis Artificial respiration should be kept up very steadily for at least three hours

Foreign Pharmacopœias — Official in Jap, Mex, Swiss and U S Not in the others

Descriptive Notes — Gelsemium Root consists mostly of the underground stem or rhizome, with occasional pieces of the root The rhizome is easily distinguished by the presence of a small, usually dark, pith, it has a purplish-brown longitudinally-fissured bark, which is thin (about 1 mm *U S P*) and shows when fractured a few silky fibres The root is yellowish-brown and tortuous, but has no pith, both root and stem have a radiate woody structure with numerous medullary rays, the bark has a bitter taste and a faint, slightly aromatic, odour The pieces vary in diameter from $\frac{1}{4}$ to $\frac{3}{4}$ of an inch (6 to 18 mm), and about 6 to 8 inches (20 or even 30 cm

U S P) in length According to Sayre, the root contains less of the active principle than the rhizome, but it resides almost entirely in the bark, and the tincture is therefore likely to vary in strength according to the proportion of bark present, it also varies in different samples, and a tincture made from the fresh rhizome is more active as a heart depressant. Under the microscope the structure of the root is remarkable for the thick medullary rays, which are about 6 to 8 cells in thickness, the cell walls being thick and pitted, but as they approach the cortical zone, the cells become larger, thinner walled, and many of the cells contain octahedral prisms of Calcium Oxalate. The cortical parenchyma has no stone cells nor laticiferous vessels, the liber has no lignified fibres, and the numerous vessels in the wood are isolated, not in groups.

Tests—Although numerous processes have been published from time to time for the assay of the preparations of Gelsemium, very few give accurate or uniform results, and those which yield uniform results are too complicated for ordinary usage. A process which gives very satisfactory and accurate results, and which, when tried in the author's laboratory, was found to justify the claims made for it, is recorded, *Proc Amer Pharm Assoc* lv. (1907), 357. As carried out on the fluid extract, the details are as follows—A measured quantity of 15 cc is evaporated at 60° C (140° F) to a soft extract, or sufficiently to dissipate the Alcohol. A measured quantity of 5 cc of Normal Volumetric Sulphuric Acid Solution, which has been previously diluted with an equal volume of Water, is added and the resulting mass allowed to disintegrate, when this is accomplished it is transferred to a 15 cc graduated cylinder and diluted to 15 cc, it is thoroughly mixed, the precipitate allowed to settle and a measured quantity of 10 cc is filtered or decanted off into a separate bottle the acid solution is washed with Chloroform, using three separate portions each of 10 cc, 5 cc and 5 cc. The chloroformic washings are in each instance separated, mixed and in turn washed with about 5 cc of slightly acidulated Water, the acid aqueous washings being mixed with the main acid solution. The mixture is rendered alkaline with Ammonia Solution and the liberated alkaloids are shaken out with three successive quantities each of 15 cc, 10 cc and 5 cc of Chloroform. A further quantity of 10 cc of Chloroform may occasionally be necessary to extract the whole of the alkaloids, their complete extraction may be determined by allowing a few drops of the chloroformic solution to evaporate, acidifying with dilute Sulphuric Acid and testing with a drop or two of Potassio-mercuric Iodide (Mayer's) Solution. The chloroformic liquids are in each instance separated, mixed, transferred to a tared flask, the Chloroform is evaporated and the residue dried until constant in weight, the weight of alkaloids multiplied by 20 and the product divided by 3 yields the percentage w/v of Chloroform-soluble Gelsemium alkaloids present in the specimen operated upon. The alkaloidal residues remaining from the above process were of a bright yellow colour, and were pure products. A sample of fluid extract

the author's laboratory in 1885 by the then official process of the *USP*, when recently examined gave the following figures specific gravity, 0.865, total solids, 8.74 p c w/v, Absolute Alcohol, 81.11 p c w/v, and when assayed according to the process described above yielded 0.37 p c w/v of Chloroform-soluble Gelsemium alkaloids.

The following constituents of Gelsemium have been described

Gelsemin — A name given to a resinoid and eclectic remedy, resembling the alcoholic extract

Dose — $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Gelsemine, Gelsemine — The crystallisable alkaloid forming crystalline salts, described by Gerrard (*PJ* (3) xiii 641) as having the formula $C_{22}H_{26}N_2O_4$, eq 405.24 and the melting point $45^\circ C$ ($113^\circ F$). Spigel says that experiments intended to establish the formula for Gelsemine (known in Germany as **Gelseminine**) as between $C_{22}H_{26}N_2O_4$ (Gerrard) and $C_{22}H_{26}N_2O_5$ have not led to a decisive conclusion, yet the results of analyses agree more closely with the latter formula.

A brittle transparent solid, crystallising with difficulty from Alcohol. It is only sparingly soluble in Water, more readily in Alcohol (90 p c), and readily in Ether and Chloroform. It dissolves in strong Nitric Acid with little or no colour. When the liquid is allowed to evaporate spontaneously in porcelain, a permanent bluish green colour is obtained. The pure alkaloid dissolves without change of colour in concentrated Sulphuric Acid, even on warming, but if not perfectly pure, a reddish or brownish colour is obtained, which gradually becomes pinkish, and on heating becomes chocolate or purple. When treated with strong Sulphuric Acid and an oxidising agent, e.g., Potassium Bichromate, a fine reddish purple or cherry red coloration is produced, rapidly changing to a bluish green or blue tint.

Dose — $\frac{1}{120}$ to $\frac{1}{2}$ grain = 0.0005 to 0.002 gramme

Care must be taken to ascertain the intention of the prescriber when any doubt exists as to whether the alkaloid or resinoid is required.

When quite free from Gelseminine, with which all early specimens were probably mixed, Gelsemine is stated (*P* 11 38) to be without action on mammals, even when injected intravenously up to $\frac{1}{2}$ gramme = $7\frac{1}{2}$ grains. Gelseminine, on the other hand, is intensely poisonous, causing a descending paralysis of the central nervous system, $\frac{1}{2}$ grain = 0.032 gramme being the calculated lethal dose for an adult. Applied locally it produces dilatation of the pupil, and it is to the action of this alkaloid, modified by the various acid resins, that the action of Gelsemium Tincture is mainly due.

Gelseminæ Hydrochloridum — Colourless crystals, soluble in Water. Known in Germany as **Gelsemininum Hydrochloricum Cryst**.

Dose — $\frac{1}{120}$ to $\frac{1}{32}$ grain = 0.0005 to 0.002 gramme

Gelseminine — A white amorphous powder, which softens at $105^\circ C$ ($221^\circ F$), and melts at $120^\circ C$ ($248^\circ F$) with partial decomposition. Insoluble in Water, soluble in Alcohol (90 p c), Ether and in Chloroform. With dilute Nitric Acid it yields a brown coloration, with concentrated Nitric Acid a green coloration, and with Sulphuric Acid a yellow coloration, changing on the addition of an oxidising agent, e.g., Potassium Bichromate, to violet and, finally, green. It is intensely bitter and poisonous.

Gelsemic Acid is not known to have any medicinal properties, but affords reactions, which to some extent serve as a test for Gelsemium preparations, particularly the blue fluorescence which it produces in alkaline solutions.

Colourless, odourless and nearly tasteless groups or tufts of prisms, or in minute scales and plates. It is not distinctly acid to Litmus paper. It is soluble in hot Water, readily soluble in Alcohol (90 p c), and in Ether and Chloroform. It dissolves in solutions of the fixed alkalis and in Ammonia.

solution, forming a solution having an intensely yellow colour by transmitted light, but which by reflected light exhibits a strong green fluorescence, which is readily destroyed by free acids. It dissolves in Nitric Acid with the production of a yellow or orange colour changing, on the addition of an excess of Ammonia solution, to a blood red coloration.

Preparation

TINCTURA GELSEMI. TINCTURE OF GELSEMIUM

1 of Gelsemium Root, in No 40 powder, percolated with Alcohol (60 p c) to yield 10 (1 in 10)

Dose.—5 to 15 minims = 0.3 to 0.9 c c

Swiss, maximum single dose, 1 gramme, maximum daily dose, 5 grammes

Foreign Pharmacopœias.—Official in Mex, 1 in 5, Swiss and US, 1 in 10, Jap, 1 in 8. All by weight except US. Not in the others.

A girl 9 years old was killed in two hours by 2 fl drim = 7.1 c c of the Tincture.

Tests.—Tincture of Gelsemium has a sp gr of 0.920 to 0.925, it contains about 1.5 p c w/v of total solids and about 58 p c w/v of Absolute Alcohol. When assayed according to the process described under Gelsemium Radix the B P Tincture yielded 0.043 p c w/v of Chloroform-soluble Gelsemium alkaloids. A specimen of the US P Tincture prepared and examined in the author's laboratory had a specific gravity of 0.913, it contained 1.8 p c w/v of total solids. When assayed according to the process recommended under Gelsemium Radix it yielded 0.048 p c w/v of Chloroform-soluble Gelsemium alkaloids.

A standard of 0.025 p c w/v of Gelsemine has been suggested for the Tincture, but a standard of total alkaloids is suggested as a safer basis.—Y P B '04, 279.

The Rhizome contains from 0.38 to 0.7 p c of total alkaloids, so that 0.5 p c might be regarded as a suitable standard, equal to 0.05 p c of total alkaloids for the Tincture.

The percentage of alkaloids in the Tincture may vary between 0.002 and 0.076 p c w/v, but standardisation, according to total alkaloids without the ratio of the two alkaloids, may be of doubtful value.

Not Official

EXTRACTUM GELSEMI ALCOHOLICUM.—Gelsemium in No 60 powder percolated with Rectified Spirit and evaporated to an extract.—B P '85.

Dose.— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme.

This has been incorporated in the B P C.

FLUIDEXTRACTUM GELSEMI.—Percolate 100 of Gelsemium in No 60 powder with Alcohol (95 p c) until exhausted, reserve the first 90 of percolate and evaporate the remainder to a soft extract, which mix with the reserved portion and make up to 100.—US P.

Average dose.—One minim = 0.06 c c.

This has been incorporated in the B P C.

GENTIANÆ RADIX.

GENTIAN ROOT

FR, RACINE DE GENTIANE, GER, ENZIANWURZEL, ITAL, GENZIANA,
SPAN, GENCIANA

The dried Rhizome and Roots of *Gentiana lutea*, L

Collected in the mountainous districts of central and southern Europe

The active principle *Gentiopicroin* is a neutral crystalline body, soluble in Water and diluted Alcohol, insoluble in Ether

Medicinal Properties—Bitter tonic, used in cases of atonic dyspepsia, the infusion is recommended in the vomiting of pregnancy, along with a mineral acid, or when a general tonic is required, as in convalescence from acute diseases or in nervous debility

For the ordinary phthisical patient nothing is better in the way of drugs to promote appetite and aid digestion than the time honoured *Mistura Gentianæ Alkalina* of the Brompton Hospital Pharmacopœia—*L* '04, 11 1827

The extract has been largely used as an excipient to form powders into pills

Official Preparations—*Extractum Gentianæ*, *Infusum Gentianæ Compositum*, and *Tinctura Gentianæ Composita*

Not Official—*Extractum Gentianæ*, *Fluidextractum Gentianæ*, *Infusum Gentianæ Compositum Concentratum*, *Aromatic Infusion of Gentian*, *Mistura Gentianæ*, *Mistura Gentianæ Alkalina*, *Mistura Gentianæ cum Soda*, *Mistura Gentianæ Acida*, *Tinctura Gentianæ*

Incompatibles—Ferrous Sulphate, Silver Nitrate, and Lead salts

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital (*Genziana*), Jap, Mex (*Genciana*), Norw, Port, Russ Span, Swed, Swiss and U S

Descriptive Notes—The official Gentian Root is that of *Gentiana lutea*. As met with in commerce, Gentian Root consists of more or less rootstock continuous with the root, the rootstock being marked with crowded rings of leaf scars, but the root is longitudinally wrinkled. It varies much in length and thickness, seldom exceeding an inch ($2\frac{1}{2}$ cm) in diameter, *BP*, 5 to 35 mm ($\frac{1}{4}$ to $1\frac{1}{2}$ inch), *USP*, Gentian Root is usually somewhat flexible and tough, with a soft fracture showing no woody tissue or medullary rays, but when recently dried is harder and brittle. Externally it is of a yellowish-brown colour, but internally more of an orange tint or reddish-yellow. The reddish colour, which causes it to be distinguished on the Continent as Red Gentian Root, is partly the result of fermentation before the root is dried, by which the characteristic odour is also more developed. The taste is sweet at first, but soon afterwards bitter. There is occasionally met with in English commerce a root with a paler fracture, known as White Gentian, which is disagreeably bitter, and should therefore not be substituted for the Official kind. It is imported from Bordeaux, and is probably derived from *Gentiana Burseri*, Lapeyr, and is not fermented before drying. Under the microscope the tissue is seen to be devoid of sclerenchymatous cells, contains minute Calcium Oxalate crystals, and rarely a few simple starch grains. The wood possesses sieve-tubes besides reticulated

solution, forming a solution turning brown on exposure to light, but which is reflected light colorless. It is readily destroyed by free acid. It dissolves in Nitric Acid with the production of an orange colour, changing, on the addition of an excess of Ammonia to a blood red coloration.

Preparation

TINCTURA GELSEMII TINCTURE OF GELSEMIUM

1 of Gelsemium Root, in No 40 powder, percolated with Alcohol (60 p.c.) to yield 10 (1 in 10)

Dose.—5 to 15 minims = 0.3 to 0.9 c.c.

Swiss, maximum single dose, 1 gramme, maximum daily dose, 5 grammes

Foreign Pharmacopœias—Official in Mex., 1 in 5, Swiss and U.S., 1 in 10, Jap., 1 in 5. Not in the others.

A girl 9 years old was killed in two hours by 2 fl. dr. = 71 c.c. of the Tincture.

Tests.—Tincture of Gelsemium has a sp. gr. of 0.920 to 0.925, it contains about 1.5 p.c. w/v of total solids and about 58 p.c. w/v of Absolute Alcohol. When assayed by the process described under Gelsemium Radix the B.P. Tincture yielded 0.043 p.c. w/v of Chloroform-soluble Gelsemium alkaloids. A specimen of the U.S.P. Tincture prepared and examined in the author's laboratory had a specific gravity of 0.913, it contained 1.8 p.c. w/v of total solids. When assayed according to the process recommended under Gelsemium Radix it yielded 0.048 p.c. w/v of Chloroform-soluble Gelsemium alkaloids.

A standard of 0.025 p.c. w/v of Gelsemine has been proposed for the Tincture, but a standard of total alkaloids is not yet established. A later paper—J. P. B. '04, 279.

The Rhizome contains from 0.38 to 0.7 p.c. of total alkaloids, so that 0.5 p.c. might be regarded as a suitable standard, equal to 0.05 p.c. of total alkaloids for the Tincture.

The percentage of alkaloids in the Tincture may vary between 0.02 and 0.076 p.c. w/v, but standardisation, based on total alkaloids without the ratio of the two alkaloids, may be of doubtful value.

Not Official.

EXTRACTUM GELSEMII ALCOHOLICUM—Gelsemium in No 60 powder percolated with Rectified Spirit and evaporated to an extract—B.P. '85

Dose— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

This has been incorporated in the B.P.C.

FLUIDEXTRACTUM GELSEMII—Percolate 100 of Gelsemium in No 60 powder with Alcohol (95 p.c.) until exhausted, reserve the first 90 of percolate and evaporate the remainder to a soft extract, which mix with the reserved portion and make up to 100—U.S.P.

Average dose—One minim = 0.06 c.c.

This has been incorporated in the B.P.C.

GENTIANÆ RADIX.

GENTIAN ROOT

FR, RACINE DE GENTIANE, GER, ENZIANWURZEL, ITAL, GENZIANA,
SPAN, GENCIANA

The dried Rhizome and Roots of *Gentiana lutea*, L

Collected in the mountainous districts of central and southern Europe

The active principle **Gentiopicroin** is a neutral crystalline body, soluble in Water and diluted Alcohol, insoluble in Ether

Medicinal Properties.—Bitter tonic, used in cases of atonic dyspepsia, the infusion is recommended in the vomiting of pregnancy, along with a mineral acid, or when a general tonic is required, as in convalescence from acute diseases or in nervous debility

For the ordinary phthisical patient nothing is better in the way of drugs to promote appetite and aid digestion than the time-honoured Mistura Gentianæ Alkalina of the Brompton Hospital Pharmacopœia —L '04, 11 1827

The extract has been largely used as an excipient to form powders into pills

Official Preparations—Extractum Gentianæ, Infusum Gentianæ Compositum, and Tinctura Gentianæ Composita

Not Official—Extractum Gentianæ, Fluidextractum Gentianæ, Infusum Gentianæ Compositum Concentratum, Aromatic Infusion of Gentian, Mistura Gentianæ, Mistura Gentianæ Alkalina, Mistura Gentianæ cum Soda, Mistura Gentianæ Acida, Tinctura Gentianæ

Incompatibles—Ferrous Sulphate, Silver Nitrate, and Lead salts

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, F₁, Ger, Hung, Ital (Genziana), Jap, Mex (Genciana), Norw, Port, Russ, Span, Swed, Swiss and U S

Descriptive Notes—The official Gentian Root is that of *Gentiana lutea*. As met with in commerce, Gentian Root consists of more or less rootstock continuous with the root, the rootstock being marked with crowded rings of leaf scars, but the root is longitudinally wrinkled. It varies much in length and thickness, seldom exceeding an inch ($2\frac{1}{2}$ cm) in diameter, B P, 5 to 35 mm ($\frac{1}{4}$ to $1\frac{1}{2}$ inch), U S P, Gentian Root is usually somewhat flexible and tough, with a soft fracture showing no woody tissue or medullary rays, but when recently dried is harder and brittle. Externally it is of a yellowish-brown colour, but internally more of an orange tint or reddish-yellow. The reddish colour, which causes it to be distinguished on the Continent as Red Gentian Root, is partly the result of fermentation before the root is dried, by which the characteristic odour is also more developed. The taste is sweet at first, but soon afterwards bitter. There is occasionally met with in English commerce a root with a paler fracture, known as White Gentian, which is disagreeably bitter, and should therefore not be substituted for the Official kind. It is imported from Bordeaux, and is probably derived from *Gentiana Burseri*, Lapeyr, and is not fermented before drying. Under the microscope the tissue is seen to be devoid of sclerenchymatous cells, contains minute Calcium Oxalate crystals, and rarely a few simple starch grains. The wood possesses sieve-tubes besides reticulated

vessels, *P G* and *Jap*. The *B P* requires that it should not yield any definite reactions with the tests for Starch. It has been found to be adulterated with 20 p c of ground olive stones, and a conviction obtained, *P J* (4) Nov 339. The *P G* permits the use of other species besides *G. lutea*, L., including *G. Pannonica*, Scop., *G. purpurea*, L., and *G. punctata*, L. The root of *Gentiana purpurea* has a branched appearance at the apex, due to several stems arising from the crown of one root, but is even more bitter than that of *G. lutea*. The roots of *G. punctata* have a similar appearance, but are a brighter reddish-brown internally. That of *G. Pannonica* is more slender, rarely exceeding 10 mm in diameter, and has more slender branches. It is likely to occur in root imported from Austria, since it occurs abundantly in the Austrian Alps, where *G. lutea* does not occur.

Tests—Gentian Root yields about 4 p c of ash. Samples examined in the author's laboratory gave from 2.2 to 5.6 p c, with an average of 3.5 p c.

Preparations

EXTRACTUM GENTIANÆ. EXTRACT OF GENTIAN

An aqueous Extract of Gentian Root, made by maceration with cold Water for 2 hours, boiling for 15 minutes, and evaporation of the strained liquid.

Dose.—2 to 8 grams = 0.13 to 0.52 gramme

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ital., Jap., Mex., Port., Russ. Span., Swed. and U.S., with cold Water, Hung., with hot Water, Ger., Norw. and Swiss, with cold Water, and purified with Alcohol, Dan. and U.S., also **Fluid Extract**, 1 in 1.

INFUSUM GENTIANÆ COMPOSITUM. COMPOUND INFUSION OF GENTIAN

Gentian Root, $\frac{1}{4}$, Dried Bitter-Orange Peel, $\frac{1}{4}$, Fresh Lemon Peel, $\frac{1}{2}$, boiling Distilled Water, 20. Infuse 15 minutes. (1 in 80)

Dose.— $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 cc

Foreign Pharmacopœias—Official in Fr. (Tisane), Gentian Root 1, cold Water 200, Swed., similar to Brit. Not in the others.

TINCTURA GENTIANÆ COMPOSITA. COMPOUND TINCTURE OF GENTIAN

Gentian Root, 2, Dried Bitter-Orange Peel, $\frac{3}{4}$, Cardamom Seeds, $\frac{1}{2}$, macerated with 20 of Alcohol (45 p c). (1 in 10)

Dose.— $\frac{1}{2}$ to 1 fl. dram = 1.8 to 3.6 cc

Tests.—Compound Tincture of Gentian has a sp. gr. of 0.965 to 0.970, it contains about 5 p c w/v of total solids and about 43 p c w/v of Absolute Alcohol.

Foreign Pharmacopœias—Official in Jap., Mex. and U.S., similar to Brit., Port., twice as strong as Brit. Not in the others.

Tinctura Gentianæ Composita—Gentian, 10, Bitter-Orange Peel, 4, Cardamom, 1, Alcohol (95 p c), 60 and 40 of Water (mixed). Percolate slowly until exhausted and make up with menstruum to 100.—U.S.P.

A simple tincture is official in most Foreign Pharmacopœias, see below.

Not Official

EXTRACTUM GENTIANÆ (U S P)—Macerate 100 of Gentian in No 20 powder with 40 of cold Water for 24 hours, exhaust by percolation with more Water, reduce the liquid to three fourths of its bulk by boiling, strain, then by means of a water bath evaporate to a pilular consistence

FLUIDEXTRACTUM GENTIANÆ (U S P)—Exhaust 100 of Gentian in No 30 powder with Alcohol (49 p c), reserve the first 80 of percolate, and evaporate the remainder to a soft extract, which dissolve in the reserved portion, and make up with Alcohol (49 p c) to 100

INFUSUM GENTIANÆ COMPOSITUM CONCENTRATUM—Gentian Root in No 10 powder, 10, Dried Bitter Orange Peel in No 10 powder, 10, Tincture of Lemon, 10, Tincture of Orange, 5, Alcohol (90 p c), 17 5, Dilute Chloroform Water (1 in 1000) sufficient to make 100 Mix the tinctures with the Alcohol, and re-percolate the drugs with dilute Chloroform Water, adding the mixed tinctures to the reserved portion

Dose— $\frac{1}{2}$ to 1 fl dm —*Fan* and *Wright*, *P J* '06, 1 165 and '07, 1 622, *C D* '06, 1 252, *Y B P* '07, 250

This appears in the *B P C*

MISTURA GENTIANÆ—Gentian Root, sliced, $\frac{1}{4}$ oz Bitter Orange Peel cut small, 30 grains, Coriander Fruit, bruised, 30 grains, Proof Spirit, 2 fl oz. Distilled Water, 8 fl oz —*B P* '87

Macerate the ingredients first in the Proof Spirit for two hours, then add the Water, macerate again for two hours, and strain through calico

Dose— $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

Gentian Root, sliced, 2 50, Bitter Orange Peel, cut small, 0 75, Coriander Fruit, bruised, 0 75, Alcohol (60 p c), 20, Distilled Water, 100 —*B P C*

Same directions as above

Mistura Amaro alkalina (Gentian Mixture) is official in Dan

MISTURA GENTIANÆ ALKALINA—Sodium Bicarbonate, 15 grains, Diluted Hydrocyanic Acid, 3 minims, Aromatic Infusion of Gentian, to 1 fl oz —*Brompton*

Aromatic Infusion of Gentian (*Brompton*)—Gentian, 2 oz, Lemon Peel, 6 dm, Orange Peel, 3 dm, Boiling Water, 1 gallon

MISTURA GENTIANÆ CUM SODA—Sodium Bicarbonate, 15 grains, Compound Infusion of Gentian, to 1 fl oz —*St Thomas's*

This has been incorporated in the *B P C*

MISTURA GENTIANÆ ACIDA—Diluted Nitro hydrochloric Acid, 10 minims, Spirit of Chloroform, 5 minims, Compound Infusion of Gentian, to 1 fl oz —*Royal Free*

Diluted Nitro-Hydrochloric Acid, 10 minims, Spirit of Chloroform, 10 minims, Compound Infusion of Gentian, to 1 fl oz —*B P C*

TINCTURA GENTIANÆ (Ger)—Gentian Root, 1, Alcohol (90 p c), 5, by weight

This is also official in Austr, Belg, Dan, Dutch, Fl, Ger, Ital, Jap, Mex, Norw, Port, Russ, Span and Swiss, 1 in 5 All by weight

Gentian Root, 1, Alcohol (45 p c), 10, by maceration —(*B P C*)

Not Official

LIQUID GLUCOSE

As met with in commerce, it is clear, almost colourless, devoid of smell, and resembles in consistence Canada Balsam It should be free from Arsenic

In exhausting diseases, subcutaneous injection of 25 grammes in 24 hours, of a 5 p c solution —*B M J* '02, 1 770

It forms an excellent excipient for pills, more particularly when diluted with Syrup

DILUTED GLUCOSE—Glucose, 3 oz, Syrup, 1 fl oz, mix.

A good excipient for pills
The following is introduced for this purpose, does not
answer so well, as it is adhesive

Official Preparation

SYRUPUS GLUCOSI. SYRUP OF GLUCOSE

Liquid Glucose, 1, Syrup, 2 Mix at a gentle heat

GLUSIDUM.

GLUSIDE

$C_7H_5NSO_3$, eq 181.77

BENZOYL SULPHONIMIDE

B P Syn—GLUCUSIMIDE

FR, SACCHARINE, GER, BENZOESAURESULFINID, ITAL, SACCARINA,
SPAN, SACARINA

A white crystalline powder, possessing an exceedingly sweet characteristic taste

Gluside is the anhydride of Ortho-sulphamide-benzoic Acid

Although the *B P* formula $C_7H_5COSO_2NH$ is attached to the synonym Benzoyl sulphonimide, it is not to be inferred that, commercially, Saccharin is sufficiently pure to allow of its representation by this or any other formula

Commercial Saccharin is not a pure product, but is 'standardised' to 300 times the sweetening power of Cane Sugar, the pure chemical (Saccharin puriss) to 500 times its weight of Sugar. The proportion of impurity may be estimated by treatment with Acetone, in which the pure salt is completely soluble

Orthobenzoisulphimide (commercial Saccharin) is put on the market as a white micro-crystalline powder containing a considerable proportion of Para-sulphamnebenzoic Acid

Commonly known as 'Saccharin'

The Saccharin Commission supplies Saccharin of the following strengths —
330, 450, 500, 550 The first corresponds with Glusidum, *B P*

Solubility—1 in 400 of cold Water, 1 in 28 of boiling Water, 1 in 38 of Alcohol (90 p c), 1 in 100 of Ether, 1 in 500 of Chloroform, 1 in 48 of Glycerin

It is also readily soluble in all alkaline solutions, either of Hydroxide, Carbonate, or Bicarbonate. It is the part of an acid and displacing Carbonic Acid. Soluble Saccharin'

Medicinal Properties—It is used as a substitute for Sugar in diabetes and hepatic diseases and to cover the taste of nauseous drugs. It is eliminated in the urine and saliva

1 grain sweetens 6 to 8 oz of fluid

Dose— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Not Official—Saccharinum Solubile, F. v. Glusid and Tabelæ Saccharini (Saccharin Discs), Sacrol

Foreign Pharmacopœias—Official in Austr, Belg, Dan., Dutch, Fr, Ital, Jap, Norw (Saccharinum), Mex (Sacarina), Russ, Span, Swed, Swiss and U S (Benzosulphimide) Not in the others

Tests—The m p of pure Benzoyl sulphimide is 224°C ($435\ 2^{\circ}\text{F}$), $223\ 5^{\circ}\text{C}$ ($434\ 3^{\circ}\text{F}$) is given in the *Fr Codex*. The *BP* m p for the substance recrystallised from warm Water is between 218° and 219°C ($424\ 4^{\circ}$ and $426\ 2^{\circ}\text{F}$), the *USP* gives 219° to 220°C ($426\ 2^{\circ}$ to 428°F). It possesses an intensely sweet taste, which is perceptible in solutions up to 1 in 100,000 of Water. When moistened with an excess of Potassium Hydroxide Solution, dried, the residue gently fused for several minutes, cooled, dissolved in Water and the solution is faintly acidified with diluted Hydrochloric Acid, it yields with Ferric Chloride T S a purple violet coloration. When fused with a mixture of Potassium and Sodium Carbonates and Potassium Nitrate, the residue dissolved in Water, and the solution filtered, the filtrate yields with Barium Chloride Solution, after acidification with diluted Hydrochloric Acid, a copious white precipitate insoluble in Hydrochloric Acid. 0.01 gramme, heated with an equal weight of Resorcin and a few drops of Sulphuric Acid, yields a mixture at first yellowish red and then greenish-brown. If the residue is dissolved in cold Water and an excess of Sodium Hydroxide (15 p c) added, the mixture assumes a strong green fluorescence. It dissolves with effervescence in warm Sodium Bicarbonate Solution forming 'soluble Gluside' or 'soluble Saccharin,' 100 parts of Gluside yielding nearly 113 parts of neutral 'soluble Gluside.'

The more generally occurring impurities are organic impurities, readily charred by Sulphuric Acid, Glucose, and other reducing Sugars, e.g. Milk Sugar, Ammonium salts, Benzoic or Salicylic Acid, and inorganic impurities. The *BP* is content with a Sulphuric Acid test for Sugar, and does not include tests, other than the m p for any of the remaining impurities. It states that no blackening should occur even when the mixture is gently warmed with Sulphuric Acid. The *USP* gives the respective quantities of substance and reagent to be employed, the temperature at which the mixture is to be maintained, and the time allowed for the test. Glucose may be detected by Potassium Hydroxide Solution, a test for Milk Sugar and a supplementary test for Glucose is afforded by Potassium cupric Tartrate (Fehling's) Solution, the substance should not evolve an odour of Ammonia when warmed with Calcined Magnesia and Water, Benzoic or Salicylic Acid are detected in a saturated aqueous solution by the Ferric Chloride test described below, and inorganic impurities by the ash left on ignition, which should amount at the highest to 0.5 p c.

Alkali Hydroxide and Sodium Bicarbonate—Gluside is readily soluble in T S of Ammonia, *BP* and *USP*, in Alkali Hydroxide solutions, *USP*, in T S of Sodium Bicarbonate with evolution of Carbon Dioxide, *BP*, and *USP*.

Sulphuric Acid—If 0.2 gramme of Benzosulphimide be dissolved with agitation in 10 c c of pure Sulphuric Acid, and the solution kept at a temperature of from 48° to 50°C ($118\ 4^{\circ}$ to $122\ 2^{\circ}\text{F}$) on a water bath, it should not within 10 minutes show a brown colour, *USP*.

Potassium Hydroxide—The solution of 0.2 gramme in 5 c c of T S of Potassium Hydroxide should be clear except after prolonged heating, *USP*.

Cupric Tartrate—A solution of Benzosulphimide in T S of Potassium

Hydroxide similar to the above should not, on heating with 5 c.c. of Volumetric solution of Alkaline Cupric Tartrate, deposit any red Cuprous Oxide, *U S P*

Ferric Chloride—No precipitate or violet colour should appear when T S of Ferric Chloride is added drop by drop to a hot aqueous solution of Benzo-sulphimide, *U S P*

Not Official

SACCHARINUM SOLUBILE ('SOLUBLE GLUSIDE')—A soluble Sodium Gluside, containing about 90 p.c. of Gluside. It is much more palatable than ordinary Gluside, which leaves a disagreeable after-taste.

This powder is soluble 1 in 15 of Water.

ELIXIR GLUSIDI *Syn* ELIXIR SACCHARINI—Dissolve 5 of Gluside with 1 of Sodium Bicarbonate in 80 of Distilled Water, add 12½ of Alcohol, filter, and wash the filter with Distilled Water to produce 100—*B P C*

This is a modification of *B P C Formulary '01*

Dose—5 to 20 minims

TABELLÆ SACCHARINI (SACCHARIN DISCS)—Contain ¼ grain = 0.032 gramme Saccharin in each. Should be readily soluble in Water, and should not contain Starch or Sugar.

Sucrol (Dulcin)—Paraphenetol Carbonide is a powerful sweetening agent which occurs in small glistening crystals, it is said to possess about 200 times the sweetening power of Sugar.

GLYCERINUM.

GLYCERIN GLYCEROL

FR, GLYCERINE OFFICINALE, GER, GLYCERIN, ITAL, GLICERINA;
SPAN, GLYCERINA

A Trihydric Alcohol, $C_3H_8O_3$, eq 91.37, containing a small percentage of Water, obtained during the saponification of fats and fixed oils by the action of alkalis, or by their hydrolysis by means of superheated steam.

Glycerin is always produced during the alcoholic fermentation of Sugar to the extent of 3 p.c. of the Sugar employed, and consequently is present in all fermented liquids.

A clear, colourless and odourless thick syrupy hygroscopic liquid, possessing a characteristically sweet taste and producing a sense of warmth in the mouth. It should be kept in well-closed vessels.

Solubility.—Mixes in all proportions with Water and Alcohol, but insoluble in Chloroform, Ether and Oils.

It possesses great powers as a solvent, and is an excellent excipient for many medicinal substances.

Medicinal Properties.—Undiluted it is an irritant, but diluted with aqueous menstrua it is emollient. It is a mild laxative. Internally it is given in irritating cough, it is recommended as a rectal injection for constipation, 1 to 2 dm, or the same diluted with an equal quantity of Water produces an evacuation very soon after the injection, also combined with Gelatin or Cocoa-nut Stearin to form a suppository for the same purpose, it is very convenient, but may aggravate hæmorrhoids if present.

Externally, a useful addition to lotions and other applications in skin diseases, as pityriasis, eczema, psoriasis, prurigo and lichen.

Used for chilblains and chapped hands, and dryness of the skin on mucous membranes, but it should be diluted with 3 parts of Water for these purposes, or applied in the form of Glycerin of Starch - Used in poultices ($\frac{1}{4}$ or $\frac{1}{8}$) it keeps them soft for a long time

It is useful in fermentative dyspepsia, when taken in 1 or 2 drm doses, and does not hinder digestion—*L* '80, ii 6, '96, ii 25

Dose—1 to 2 fl drm = 3.6 to 7.1 c.c.

Smaller doses are usually prescribed

Prescribing Notes—*It is much employed as a sweetening agent in the place of Syrup, and is better for covering the unpleasant astringent taste of Iron Perchloride, it is largely used in pharmaceutical preparations as a solvent, and, being an antiseptic, it also acts as a preservative. Mixed with equal volumes of Syrup, Alcohol and Mucilage, it forms a good pill excipient. It is too hygroscopic to be used alone.*

Official Preparation—Suppositoria Glycerini Used in the preparation of Extractum Cinchonæ Liquidum, Extractum Saisæ Liquidum, of all the Glycerina and Lamellæ, Linimentum Potassii Iodidi cum Sapone, Liquor Ethyl Nitritus, Liquor Thyroidei, Lotio Hydrargyri Nigra, Mel Boracis, Pilula Ferri, Pilula Quinina Sulphatis, Syrupus Pruni Virginianæ, Tinctura Kino, Tinctura Rhei Composita, Unguentum Acidi Carbolici, Unguentum Iodi, and Unguentum Suppuris Iodidi

Not Official—Dispensing Syrup, Glycerin with Rose Water, Suppositoria Glycerini, Suppositoria Glycerini cum Stearino

Foreign Pharmacopœias—Official in Austri, sp gr 1.250, Belg, sp gr. 1.240, Dan, Ger, Hung, Jap, Norw, Russ and Swed, sp gr 1.225 to 1.235, Dutch, sp gr 1.230 to 1.235, Fr, sp gr 1.264, Ital, sp gr 1.226 to 1.260, Mex, Port and Span, sp gr 1.260, Swiss, sp gr 1.224 to 1.235, U.S., not less than 1.246 at 25° C (77° F)

Tests—Glycerin has a sp gr of 1.260, which figure is given in the *BP*, the *USP* gives not less than 1.246 at 25° C (77° F.), the *PG* 1.225 to 1.235. The aqueous solution is neutral to Litmus paper. When heated in an open capsule it yields irritating acid vapours of Acrolein. Dilute aqueous solutions are slowly volatilised with the vapour of Water, whilst stronger solutions rapidly volatilise at boiling temperatures. A loop of Platinum Wire, containing a fused bead of Borax moistened with Glycerin, imparts to the edge of a non-luminous flame a transient vivid green colour. When boiled with Potassium or Sodium Hydroxide and Potassium Permanganate Solution, the latter is immediately reduced. The filtered liquid, when made faintly acid with Acetic Acid, yields with Calcium Chloride Solution a white precipitate, insoluble in Acetic Acid, soluble in Hydrochloric Acid. This reaction with alkaline Permanganate forms the basis of a method for the determination of Glycerin which, in the absence of foreign bodies yielding Oxalic Acid on oxidation, has been proved to give very accurate results.

The more generally occurring impurities are those of an inorganic nature, *e.g.*, Arsenic, Copper, Lead, Iron, Calcium, Potassium, Sodium, Ammonium, Chlorides and Sulphates, and mineral impurities, those of an organic nature, Sugars, *e.g.*, Grape and Cane Sugar, foreign organic matter, *e.g.*, Acrolein, Formic Acid or Formates, Butyric Acid, Oxalic Acid or Oxalates, and organic impurities readily charred by Sulphuric Acid. The *BP* employs Siebold's modification of the

Gutzeit's test for the detection of Arsenic, which approximately indicates 1 part of Arsenic in 250,000 of Glycerin. The *P G* employs the Bettendorf's test, the *U S P* then modified Gutzeit's test, which indicates less than 1 in 100,000. A standard of not less than 2 parts of Arsenic per million is suggested (*C D '08*, 1796). A very great majority of about 450 samples mentioned in this reference showed less than this figure, 4 parts per million, however, may be considered by some to be a sufficiently low limit. Copper, Lead and Iron are detected by Hydrogen Sulphide Solution, the two former in slightly acid, the latter in alkaline solution, Calcium by Ammonium Oxalate Solution, Potassium and Sodium in the residual liquid after separation of the other metals, Ammonium by boiling with Potassium or Sodium Hydroxide Solution. The three latter, however, are unlikely impurities. Chlorides and Sulphates are detected by Silver Nitrate and Barium Chloride or Nitrate solutions respectively.

The *B P* and the *U S P* employ Potassio-cupric Tartrate (Fehling's) Solution as a test for Cane and Grape Sugars, the *B P* requiring that no precipitate shall be produced even after previous acidification and boiling. The *U S P* specifies quantities of substance and reagent to be used for the inversion, quantity of reagent to be used for precipitation, and time limit within which no cloudiness or precipitate is permitted. Acrolein, Formic Acid or Formates, classed by the *B P* as foreign organic matter, may be detected by Silver Ammonio-nitrate Solution. With regard to this test more explicit directions are contained in the *P G* monograph than in either the *B P* or the *U S P*, the quantities and temperature to which the mixed solutions are to be heated are given, and a definite interval of time (5 minutes), during which neither coloration nor brownish-black deposit should appear, are given. The test for Butyric Acid is, save for the difference in the strengths of the Alcohol, virtually the same in the *B P* and the *U S P*. The *U S P* and *P G* include a test for Oxalic Acid with Calcium Chloride TS, but no test appears in the *B P*. The *U S P* adopts a time limit of one hour during which, in testing for acidly charred organic impurities, the mixture of Glycerin and Sulphuric Acid is required to develop a colour not darker than yellow. The *B P* adopts no time limit. Glycerin should be entirely dissipated when heated at a high temperature, and on ignition should leave no fixed residue. Each of the above tests appears in small type below, with a further detailed comparison between the pharmacopœial methods of application.

Potassio cupric Tartrate—Even after it has been treated with mineral acid and boiled, Glycerin should give no red precipitate when mixed with excess of TS of Potassio-cupric Tartrate, *B P*, 5 c.c. of Glycerin mixed with 50 c.c. of Water and 10 drops of Hydrochloric Acid in a small flask, and heated for half an hour on a water-bath, 10 c.c. of this hot liquid mixed with 2 c.c. of Sodium Hydroxide TS and 1 c.c. of Alkaline Cupric Tartrate Volumetric Solution should show no yellowish-red cloudiness or precipitate within 6 hours, *U S P*.

Sulphuric Acid—It is officially required to yield either no coloration at all, or at the most a very pale straw coloration when 5 c.c. of Glycerin is shaken with 5 c.c. of Sulphuric Acid, care being taken to keep the mixture well cooled.

A mixture of 5 c c each of Glycerin and Sulphuric Acid should acquire, on standing for 1 hour, a colour not darker than yellow, *U S P*

Diluted Sulphuric Acid—1 c c of Glycerin, warmed gently with 1 c c diluted Sulphuric Acid, should not give off an unpleasant rancid odour, *P G*

Alcohol (90 p c) and diluted Sulphuric Acid—It is officially required that no fruity odour should be produced when equal volumes of Glycerin and a mixture of Alcohol (90 p c) and diluted Sulphuric Acid are gently heated together. 5 c c of Glycerin mixed with an equal volume of Alcohol (94.9 p c) and diluted Sulphuric Acid and gently heated, no fruity odour should be recognisable, *U S P*

Barium Nitrate or Chloride—A portion of an aqueous solution (1-5 *P G*, 1-10 *U S P*) should be unaffected by T S of Barium Nitrate *P G*, by T S of Barium Chloride, *U S P*

Ammonium Oxalate—A portion of an aqueous solution, as above, should be unaffected by T S of Ammonium Oxalate, *P G* and *U S P*

Calcium Chloride—A portion of an aqueous solution, as above, should be unaffected by T S of Calcium Chloride, *P G* and *U S P*

Silver Nitrate—A portion of an aqueous solution, as above, should yield no colour, cloudiness or precipitate with T S of Silver Nitrate, *U S P*, should yield not more than an opalescent turbidity, *P G*

Ammonia and Silver Nitrate—It is officially required that at the ordinary temperature no darkening in colour should be produced when a few drops of Silver Nitrate Solution are added to a mixture of Glycerin and Ammonia Solution in equal volumes. 1 gramme of Glycerin and 1 c c of Ammonia T S warmed on a water-bath to 60° C (140° F) and then mixed with 3 drops of Silver Nitrate T S, there should be neither coloration nor a brownish black deposit in the mixture within 5 minutes, *P G*, no colour, cloudiness or precipitate should appear when an aqueous solution of Glycerin (1-10) is treated in a test tube with Ammonio Silver Nitrate T S, the tube being loosely stoppered to protect it from impurities, and allowed to stand, protected from light, for at least 5 minutes, *U S P*

Sodium Hydroxide—1 c c of Glycerin warmed with Sodium Hydroxide T S should neither become coloured nor evolve ammonia or any odour resembling that of glue, *P G*

Stannous Chloride—A mixture of 1 c c of Glycerin and 3 c c of Stannous Chloride T S should not assume a dark colour in the course of an hour, *P G*

Hydrogen Sulphide—An aqueous solution (1-5) should not be affected by T S of Hydrogen Sulphide, *P G*, an aqueous solution (1-20), acidified with Hydrochloric Acid, should not respond to the time limit test for heavy metals, *U S P*

Gutzeit's Test—5 c c of an aqueous solution (1 in 10) should not respond to the modified Gutzeit's test for Arsenic, *U S P*. It is officially required that within 15 minutes no yellow stain should be produced on a piece of filter paper which has been previously moistened with a drop or two of Mercuric Chloride T S and dried, and which is supported over the mouth of a test-tube containing 1 gramme of Arsenic free Zinc, 5 c c of an aqueous 2.2 p c solution of Hydrochloric Acid (*B P*) and 2 c c of Glycerin.

Preparations.

SUPPOSITORIA GLYCERINI GLYCERIN SUPPOSITORIES

Using a tared basin, $\frac{1}{2}$ oz of Gelatin, cut small, is covered with Distilled Water, which after two minutes is poured away. When the Gelatin is quite soft, dissolve in $2\frac{1}{2}$ oz of Glycerin (by weight) on a water-bath, and then evaporate the excess of Water until the product weighs 1563 grams. The mass will contain about 70 p.c. w/w of

Glycerin It may be moulded into any convenient size when required

A similar preparation has been in use for many years (*Companion* 1877) as a basis for medicated Pillaries and . . . The formula in the *Companion* arrives at the same result (70 p c) *without* evaporation. It is easy by evaporation to obtain a product containing 80 p c of Glycerin. The consistency of the mass will vary somewhat with the quality of the Gelatin, see p 556

Foreign Pharmacopœias—Official in Austri, Jap and U S, Glycerin, Sodium Carbonate and Stearin, Belg, Fr, Mex, and Swiss

Glycerin Suppositories are much more convenient to use when made with Cocoa-nut Stearin, see below

Not Official

DISPENSING SYRUP—Glycerin, Syrup, Alcohol (90 p c), and Mucilage of Acacia equal volumes

An excipient for pills Glycerin by itself is too hygroscopic

GLYCERIN WITH ROSE WATER—Glycerin, 1, Rose Water, 3, mix.

SUPPOSITORIA GLYCERINI (USP)—Dissolve 1 of . . . Sodium Carbonate in 10 of Water and add to it 60 of Glycerin (t . . . 4 of Stearic Acid, heat carefully on a water-bath until effervescence ceases and the liquid is clear. This quantity is for 20 rectal suppositories, which must be kept in tightly-covered glass vessels

The *BPC* describes it as using equal parts of Monohydric and Sodium Carbonate and Water, but this is due to the omission of a decimal point

SUPPOSITORIA GLYCERINI C STEARINO—Glycerin, 20 grains, Cocoa-nut Stearin, 40 grains, melt the Stearin, and when just fluid stir in the Glycerin and continue the stirring until the mixture becomes solid. Melt the mass with the least possible heat, and pour into moulds

They can be used without any lubricant

UNGUENTUM GLYCERINI See GLYCERINUM AMYLICUM

GLYCYRRHIZÆ RADIX.

LIQUORICE ROOT

FR, RUSSESSA GLR, SSSSHOLZ, ITAL, LIQUIRIZIA, SPAN, RLGALIZ

The Root and . . . Stem, both peeled, of *Glycyrrhiza glabra*, . . .

In the *USP* the unpeeled Spanish and the peeled Russian root are both official, in the *PG* the Russian peeled root is ordered

The principle Glycyrrhizin is comparatively tasteless, the characteristic sweetness being only developed by combination with alkali. It exists in the drug as a combination with calcium

Medicinal Properties.—A demulcent and expectorant in bronchial catarrh and cough. The liquid extract helps to disguise the taste of nauseous medicines, but many persons object to the taste of Liquorice. In the form of extract and its solution it is a domestic remedy for cough. The compound powder is chiefly valuable on account of the Senna and Sulphur it contains, and is an agreeable and mild purgative, well adapted for weak persons and for cases of hæmorrhoids

Official Preparations of Liquorice—Of the Root, Extractum Glycyrrhizæ, Extractum Glycyrrhizæ Liquidum, Liquor Sarsæ Compositus Concentratus, Pilula Hydragryi, and Pulvis Glycyrrhizæ Compositus, of the **Extract**, Confectio Sennæ and Decoctum Aloes Compositum, Extractum Glycyrrhizæ Spirituosum, of the **Liquid Extract**, Mistura Sennæ Composita and Tinctura Aloes

Not Official—Elixir Adjuvans, Elixir e Succo Glycyrrhizæ seu Elixir Pectorale, Glycyrrhizinum Ammoniatum, Mistura Glycyrrhizæ Composita, Pulvis Amygdalæ Laxativus, Syrupus Glycyrrhizæ, and Trochisci Glycyrrhizæ

Foreign Pharmacopœias—Official in all the Pharmacopœias, Austr., Belg., Dan., Dutch, Fr (Régisse), Hung., Ital (Liquirizia), Jap and Ger (Liquiritia), Mex (Orozuz), Port (Alcacus), Russ., Span (Regaliz), Swiss and U.S., all *G. glabra*

Descriptive Notes—The Liquorice Root of commerce exists in several forms. The English root is never sold in the decorticated form, but either fresh or dried. Some of the fresh Liquorice Root of commerce also comes from France. The dried Liquorice Root consists of the product of at least two plants.

That derived from France, Spain and Sicily is the product of *G. glabra*, but that from Russia, Asia Minor and Persia is chiefly the product of *G. glandulifera*, and is recognisable by its redder tint, more scaly surface, and slight acidity and bitterness. As a rule the Liquorice Root of commerce consists of a larger proportion of underground stem than of root. As the root is sweeter than the stem, samples richest in root are of greater value. The root of *G. glabra* has a thin brown bark, marked here and there with short transverse scars, is yellowish within and has a radiate and porous woody structure and a fibrous fracture, a sweet taste and characteristic odour and flavour when chewed. The underground stem does not exhibit transverse scars, but at the cut ends shows a small central depression caused by shrinkage of the pith.

French Liquorice Root is usually of good quality, and is also sold in the decorticated or peeled form. Spanish Liquorice Root from Tortosa occurs in trimmed bundles of uniform size and length and of fairly uniform pieces, that from Alicante in loosely packed bales, the root being of varying size and quantity. Sicilian Liquorice is usually peeled and, like the French, often cut up into short pieces 1 inch or less in length. Russian Liquorice is sold both in the unpeeled and in the peeled state. It is often in very large tapering pieces, sometimes blackened and hollowed near the crown of the root. It gives a paler powder than the French and Sicilian, and is somewhat acid and bitter. Persian or Bussorah root is in long cylindrical pieces, an inch or more in diameter, and is not sold in the peeled state. The official root is limited to peeled root and peeled subterranean stem of *Glycyrrhiza glabra*, Linnaeus and other species. The last three words seem superfluous, since the character* given evidently excludes that of *G. glandulifera*, Waldest and Kit, which affords the Liquorice Root of Eastern Europe and Western Asia, some of the characters of the official drug being that it should be free from bitterness and that it should be dark brown in colour, longitudinally wrinkled, but not scaly, which are the distinctive characters of the root of *G. glandulifera*.

It is not stated how these characters are to be ascertained from the root which is already peeled as purchased in commerce, and directions for peeling the dried root are not given. The acidity of Eastern Liquorice Root is due to a resin contained principally in the bark, and the bitterness to a principle named Glycamarin, the sweetness is due to Glycyrrhizic Acid which exists partly in combination with Ammonia in the fresh root, in which state it is sweeter, since it is then more soluble in Water.

Tests.—The ash of Liquorice Root amounts to from 3 to 4 p c, and should not much exceed the latter figure.

Samples of fine English root examined in the author's laboratory yielded on an average 1.5 p c of ash, 1 p c of the English decorticated powdered root left from 3.4 to 3.5 p c of ash, with an average of 3.76 p c.

Preparations

EXTRACTUM GLYCYRRHIZÆ. EXACT OF LIQUORICE

In aqueous extract prepared by cold percolation, coagulation of Albumen at 212° F (100° C), and the subsequent evaporation to a soft extract.

Dose.—5 to 30 grains = 0.32 to 2 grams.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr. (Ext. Reglisse), Hung., Ital., Jap., Mex., Port., S. and Spain, Prussia, root with cold Water, U.S. (Extractum Glycyrrhizæ Purum), from root with Water and Ammonia. The Crude Extract sticks (*Succus Liquiritiæ*) is official in Austr., Dan., Dutch, Fr., Germany, Jap., Norw., Russ., Swed. and Swiss, U.S. (Extractum Glycyrrhizæ), Dratum from Crude Extract is official in Austr., Ger., Hung., Norw., and Swe.

Under the name Liquorice Juice, aqueous extract, prepared by boiling the root with Water, is commercial the form of sticks, Solazzi Juice is the best known brand.

EXTRACTUM GLYCYRRHIZÆ LIQDUM. LIQUID EXTRACT OF LIQUORICE

An aqueous fluid extract, treated thence as for the extract, but evaporated to sp. gr. 1.2, to this is added of its volume of Alcohol (90 p c).

Dose.—1 to 1 fl. dr. = 1.8 to 3 c.

The finished product is slightly acid.

Ammonia is added for preserving its sweetness; so long as the alkalinity is maintained that is no danger of the forming deposit which is often seen at the bottom of the bottle of the Extract of Ice cream.

Fluidextractum Glycyrrh. (U.S.P.)—1 in 1 Fluid extract, obtained by treating 100 of the Liquorice with boiling Water until exhaustion, evaporating the liquid to 45, and adding 45 of Alcohol (95 p c), after 3 days miter, and evaporate to 50, and Glycerin 25, Ammonia Water 5, Alcohol (95 p c) 20, and Water q.s. to make.

Foreign Pharmacopœias—Belg., Alcohol (30 p c), Mex., Ammonia and Alcohol, Swed., Ammonia and Alcohol, Swiss, with Chloroform Water and Alcohol (90 p c).

Test.—Liquid Extract of Liquorice varies very considerably in its character. A good liquid extract prepared from fresh English

root had a sp gr of 1.130 to 1.135, contained from 33.6 to 37.6 p c w/v of total solids and about 18 p c w/v of Absolute Alcohol.

The palatability of commercial liquid extracts also manifests considerable variation, some being almost bitter in taste, and few bearing any comparison with a liquid extract made from fresh English root.

EXTRACTUM GLYCYRRHIZÆ SPIRITUOSUM

Dissolve 10 of Extract of Liquorice in a small quantity of Distilled Water, add 5 of Alcohol (90 p c) and make up with Distilled Water to 20.

Dose — $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c c

This is official in the *Ind* and *Col Add* for India and the Eastern Colonies.

Extracta Liquida—Any Liquid Extract, defined in the Text of the Pharmacopœia, containing less than one fourth of its weight of Alcohol (90 p c), may have the proportion of Alcohol (90 p c) increased, to an extent not exceeding one fourth of the weight of the Extract, in India and other tropical countries where otherwise the preparation would be liable to ferment.

PULVIS GLYCYRRHIZÆ COMPOSITUS COMPOUND POWDER

OF LIQUORICE

N O Syn—PULVIS LIQUIRITIÆ COMPOSITUS, PULVIS PECTORALIS

KURELLÆ

Senna, 2, Liquorice Root, 2, Fennel Fruit, 1, Sublimed Sulphur, 1, Refined Sugar, 6

Dose — 60 to 120 grains = 4 to 8 grammes

As a mild aperient, a teaspoonful or more for adults, less in proportion for children.

For diabetic patients the late Balmanno Squire suggested that the Sugar and Liquorice should be replaced by Almond-meal and Powdered Gum Acacia. It can be ordered as *Pulvis Amygdalæ Laxativus*. See p 574.

Tests—Compound Liquorice Powder usually contains 30 to 50 p c and should not contain more than 60 p c of moisture, from the ash ranges from 4 to 6 p c, and the soluble ash from 2 to 3 p c, averaging about 2.5 p c. The percentage of Sulphur may be determined by extracting the sample with Carbon Bisulphide, evaporating off the solvent, treating with Fuming Nitric Acid, filtering, Bromine, and a few crystals of Potassium Chlorate. A little Nitric Acid by evaporating to a small bulk with Hydrochloric Acid, removing the moistening with Hydrochloric Acid, and again evaporating to a small bulk. The Sulphur is oxidised to Sulphuric Acid, which may be determined in the usual manner by precipitation with Barium Chloride Solution. It is generally present to the extent of about 8 to 9 p c. Ground Olive stones, Maize Starch, exhausted drugs, etc., Olive stones and Maize Starch may be detected by the microscopical appearance of the specimen. A criterion of exhausted drugs is afforded by a joint determination of the percentage w/v of the total extractive matter yielded to Alcohol (70 p c) and the percentage of Sugar, the difference between these determinations should be from 10.5 to 13.0 p c. A specimen of compound Liquorice powder prepared in the author's laboratory from the finest English decorticated

TROCHISCI GLYCYRRHIZÆ—Extract of Liquorice, 18, Anise Oil, 3, Acacia Lozenge Mass, 60, to make six lozenges—*Brompton*, and *Chry Chest*

This has been incorporated in *B P C* giving the synonym 'Brompton Cough Lozenge'

U S P has a *Trochiscus Glycyrrhizæ et Opii*, containing 15 grammes of Extract of Glycyrrhiza, 0.5 gramme of Powdered Opium, and 0.2 cc of Oil of Anise in 100

Fr has *Pâte de Réglisse officinale* containing about 0.02 pc of Extract of Opium

GOA POWDER *See* ARAROA

GOSSYPIUM.

COTTON

B P Syn—COTTON-WOOL

Fr, COTON HYDROPHILE, *GER*, GEREINIGTE BAUMWOLLE, *ITAL*, COTONE ASSORBENTE, *SPAN*, ALGODON HIDROFILO

The Hairs of the Seed *Gossypium Barbadosense*, *L*, and of other species of *Gossypium*, from which, by suitable treatment, the fatty matter has been removed. This is commonly known as **Absorbent Cotton-Wool**

Cotton Wool is medicated with Carbolic Acid, Salicylic Acid, Boric Acid, Eucalyptol, Thymol, Iron salts, Mercuric Chloride, Double Cyanide, Sal Alembroth, Iodine, Iodoform, and other substances

Official Preparation—Used in the preparation of Pyroxylin

Foreign Pharmacopœias—*Austr*, *Dutch*, *Ger*, *Jap*, *Russ*, *Swed*, and *Swiss* (*Gossypium Depuratum*), *Ital* (*Cotone Assorbente*), *Mex* (*Algodon* and *Algodon hydrofilo*), *Port* (*Algodoeiro*), *Span* (*Algodon*), *US* (*Gossypium Purificatum*), *Fr* (*Coton Hydrophile*), not washed, *Belg* (*Coton hydrophile*). Not in the others. *Medicated Cottons* have been inserted in *Dutch* and *Mex*

MOUTH AND NOSE PROTECTOR—For use in poisonous and injurious trades. *Squire and Sons* exhibited this respirator at the International Health Exhibition (1884) and obtained for it a bronze medal. It consists of layers of washed and sterilised Cotton-Wool placed between layers of Perforated Zinc and Perforated Cardboard, formed into a pliable respirator which covers the mouth and nose

Gamgee Tissue or Absorbent Gauze and Cotton Wool Tissue, which consists of layers of absorbent Cotton-Wool enclosed in absorbent Gauze, is a favourite dressing, and is convenient for applying lotions

Tela Depurata Purified Mull (*Ger*)—This mull should have a breadth of 100 centimetres, and each square metre should weigh at least 30 grammes, and each square centimetre should contain at least 24 threads, when not otherwise ordered

Not Official

GOSSYPH RADICIS CORTEX

The Bark of the Root *Gossypium herbaceum*, *L*, and of other species of *Gossypium*.

It is official in the *Ind* and *Col Ad* for India and the Eastern, North American and West Indian Colonies

Medicinal Properties—The Tincture and Fluid Extract have been used in America, and occasionally in Europe, as a substitute for Ergot in labour, and to check metrorrhagia—*L* '94 u 1298

Foreign Pharmacopœias—Official in U.S. Not in the others

Descriptive Notes—The bark of the root as met with in commerce occurs in the form of thin, flat or slightly curled strips more or less curled in drying, dark brown externally, with a thin outer layer which, when abraded, shows a reddish brown coloured layer beneath. The inner surface is of a yellowish-white colour when recently dried, but darkens into brownish-red when the bark is kept, it is finely striated with projecting medullary rays. The transverse section shows the bast in radiating lines which are broader at the base. The transverse fracture is laminate and fibrous, and although the bark is easily split longitudinally, it is only broken transversely with difficulty. It has very little odour and an astringent and faintly acid taste.

A spurious cotton bark is sometimes met with which has a dark brown inner surface and is more easily broken transversely.

DECOCTUM GOSSYPII RADICIS CORTICIS (*Ind and Col Add*)—Boil 4 of Cotton Root Bark with 40 of Distilled Water until reduced to 20, strain and make up to 20.

Dose— $\frac{1}{2}$ to 2 fl oz = 14.2 to 56.8 cc

For India and the Eastern, North American and West Indian Colonies

This has been incorporated in the *B P C*

EXTRACTUM GOSSYPII RADICIS CORTICIS LIQUIDUM (*Ind and Col Add*)—A 1 in 1 fluid extract of Cotton Root Bark prepared by percolation, using as a menstruum first Alcohol (90 p c), containing 25 p c Glycerin, and finally Alcohol (90 p c)

Dose—30 to 60 minims = 1.8 to 3.6 cc

For India and the Eastern, North American and West Indian Colonies

This has been incorporated in the *B P C*

TINCTURA GOSSYPII—Dried Bark of the Root of the Cotton Plant in powder, 1, percolate with sufficient Alcohol (60 p c) to produce 4.

Dose—1 fl dr = 3.6 cc

This has been incorporated in the *B P C*

GRANATI CORTEX.

POMEGRANATE BARK

FR, Ecod, RCE DE GRENOBLE, GER, GRANATRINDE, ITAL, MELOGRANATO, SPAN, CORTEZA DE GRANADO

The dried Bark of the Stem and Root of *Punica Granatum*, L

Neither the *B P* nor the *U S P* requires the Bark to yield any definite percentage of alkaloids. The *P G* requires it to yield 0.4 p c w/w of alkaloids when determined by the process given below. The *Fr Codex* (1908) requires the dried bark to yield not less than 0.25 p c of alkaloids.

The Pomegranate-root alkaloids are Pelletierine (Punicine), Isopelletierine (Isopunicine), Granatoline, Pelletierine (Methylpunicine) and Pelletierine (Methylpunicine). The first two constitute the Pelletierine of medicine, the last two are inactive. Pelletierine is a volatile liquid, but forms stable salts.

Medicinal Properties—considered effective in killings.—Astringent and anticholeric. It is ceded and followed by a purgative tapeworm; the dose should be pre- same purpose. Pelletierine Sulphate is used for

Incompatibles.—Alkalis, Limes.

Official Preparation.—Decoctum, Water, Metallic salts, Gelatin

Not Official.—Extractum Granati, Granati Corticis Tannas, Pelletierine Sulphas and Pelletierine

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Jap, Fr (Grenadier), Ger, Hung, Ital (Melogranato), Port (Romeira), Mex, Russ and Span (Granado), Swiss and U S Not in Norw or Swed

Descriptive Notes—The commercial article consists chiefly of the stem bark, which can be distinguished by the presence of lichens and of a dark green phelloderm layer from that of the root, which has conchoidal depressions and is more or less curved or twisted, that of the stem being straight. The colour is yellowish grey externally and brownish-yellow on the inner surface. The fracture is short and of a pale yellow, and presents under a lens a tessellated or latticed appearance, from the presence of numerous fine radial lines crossed by fine tangential lines. It is about $\frac{1}{12}$ in thickness. The official bark is stated to be 2 to 4 in ($\frac{1}{2}$ to 1 dm) in length, and $\frac{1}{2}$ to 1 in (12 to 25 mm) in width. The taste is very astringent, with a slight bitterness, but it has no distinctive odour. As the root bark may contain six times as much alkaloid as the stem bark, the drug is more valuable in proportion to the amount of root bark it contains.

When examined under the microscope the powdered bark should contain no other ingredients except round, single starch grains with a diameter of 0.0025–0.008 mm, rarely compound starch grains, characteristic cork cells, sclerenchymatous cells, cells containing single or clustered crystals of Oxalate, parenchymatous cells and sieve tubes.—*P G* and *Jap*

Tests—Pomegranate Bark when allowed to macerate for an hour in slightly acidified Water, yields a yellowish solution. The addition of a few drops of Ferric Chloride Test solution to a portion of this liquid affords a bluish-black coloration, which is changed to a yellowish-red on the addition of five times the volume of Lime Water and on standing an orange red flocculent precipitate is thrown off. The process for the determination of the alkaloids officially adopted by the *P G* is as follows.—A weighed quantity of 12 grammes of Pomegranate Root dried at 100° C (212° F), in a fairly fine *Grindel* is introduced into a stoppered vessel and vigorously shaken with 90 grammes of Ether and 30 grammes of Chloroform. A measured quantity of 10 cc of a mixture of 2 parts by weight of Hydroxide Solution (15 p.c w/w) and 1 part by weight of slightly acidified Water is then added, and the mixture allowed to stand with intense shaking. A measured quantity of 10 cc of Water is added to cause the powdered root to agglomerate. The mixture is shaken and the Chloroform-ether solution is removed. A weighed quantity of 100 grammes of ether solution is filtered (after an interval of 15 minutes) when vigorously shaken, a dry, well-covered filter into a separate flask. The clear Chloroform-normal Volumetric Hydrochloric Acid (10 N) is added (10 cc), through separation the latter is filtered through a filter, and the alkaloids are washed with Water into a flask of 100 cc. The filtrate is repeated with 3 separate quantities of 10 cc of Water, the aqueous shakings being filtered through a small filter paper moistened with Water and the filtrates are diluted with Water to 100 cc. The filtrate is then filtered through a filter, the latter is washed with Water and the filtrate is diluted with Water to 100 cc. The filtrate is then filtered through a filter, the latter is washed with Water and the filtrate is diluted with Water to 100 cc.

100 c c A measured quantity of 50 c c is transferred to a stoppered flask of about 200 c c capacity, and about 50 c c of Water and sufficient Ether to form a layer of about 1 cm added After the addition of 5 drops of Iodeosin Solution, Hundredth-normal Volumetric Potassium Hydroxide Solution is added until the lower aqueous layer assumes a pale rose red coloration, the mixture being well shaken after each addition, not more than 11 c c of the solution should be necessary to produce this coloration The amount of Pelletierine corresponding to this titration figure is 0.395 p c w/w Using the mean molecular weights of Pelletierine and Methyl-pelletierine in the calculation the process amounts to not less than 0.41 p c w/w The *Ph. Code* (1900) process is a volumetric one, the result of the titration being calculated from a factor based on the mean molecular weights of Pelletierine and Methyl-pelletierine The process is carried out on the bark dried at 100° C (212° F), which is required to yield not less than 0.25 p c of alkaloids The percentage of alkaloid varies between 0.5 to 0.7 p c w/w, and may even be as high as 1.0 p c w/w The percentage of ash varies between 5.0 and 13.0 p c, and should not exceed 15.0 p c

Preparation

DECOCTUM GRANATI CORTICIS. DECOCTION OF POMEGRANATE BARK

Boil 4 oz of Pomegranate Bark with 24 fl oz of Distilled Water for ten minutes, strain and wash the residue with Distilled Water, *qs* to yield 20 fl oz (1 in 5)

Dose — $\frac{1}{2}$ to 2 fl oz = 14.2 to 56.8 c c

Foreign Pharmacopoeias — Official in Belg, 1 and 6, boil to 4, Fr (Apo. me), 1, Ital, 1 in 50, Port, 1 and $7\frac{1}{2}$, boil to 5, Spain, 1 in 10 Not in the others

0

Not Official

An excellent remedy for tapeworm is as follows —

Bruised Root-bark of Pomegranate, 2 oz, Boiling Water, 24 fl oz, macerate for 24 hours, and then boil till reduced to 18 fl oz A third part early in the morning, a third part again in half an hour, and the remainder in another half-hour A dose of Castor Oil should have been taken the previous morning, and solid food abstained from on that day This rarely fails to bring away the entire worm in two hours, and the head (at the thinnest end) should be diligently sought for This form was given in *Companion* 1873

EXTRACTUM GRANATI — Exhaust Pomegranate Root-bark with Alcohol (60 p c), distil off the Alcohol and evaporate to the consistence of an Extract 10 of Root-bark yield 34 of Extract

Foreign Pharmacopoeias — Official in Austr, Dutch, Hung, Port and Russ Not in U. S.

Fluidextractum Granati is official in U. S., is prepared by percolating 100 grammes of Pomegranate in No 80 powder with a mixture of Glycerin 10 c c and Alcohol (49 p c) *qs* to make 100 c c of Fluidextract

PELLETIERINA Pelletierine, $C_8H_{15}NO$ eq 140.10 — A colourless, oily liquid having an aromatic odour, and becoming brown on exposure to the air It should be kept in well stoppered glass bottles in a dark amber tint and in a cool atmosphere

Tests—Pelletierine has a sp gr at 0° C (32° F) of 0.988, and a boiling point of 195° C (383° F), at which temperature it distils.

It is soluble in Water and readily soluble in Alcohol (90 p c), Ether and Chloroform.

PELLETIERINÆ SULPHAS—A white, crystalline, non hygroscopic mass, which should be preserved from the light. It is official in *Fr. Codex* (1908).

Dose—6 grains = 0.4 gramme, prescribed with 7 grains = 0.46 gramme of Tannic Acid.

The **Pelletierine de Tanret** has been improperly called Pelletierine Tannate, on account of its being a mixture of Pelletierine Sulphate and Tannin, but it is quite distinct from the true Tannate.

PELLETIERINÆ TANNAS Pelletierine Tannate, Punicine Tannate—A yellowish, amorphous, odourless powder, prepared from Pomegranate Bark. It possesses an astringent taste.

Solubility—1 in about 700 of Water, 1 in 80 of Alcohol.

Dose—5 to 8 grains = 0.32 to 0.52 gramme.

Foreign Pharmacopœias—Official in Ital and U.S.

Tests—Pelletierine Tannate, dried over Sulphuric Acid and heated, turns brown at 20° C (302° F) and softens at about 165° C (329° F). It is faintly acid in reaction towards blue Litmus paper. The aqueous solution affords a precipitate with Mercuric potassium Iodide (Mayer's) Solution, the precipitate becoming granular and yellow coloured. It yields a white precipitate with Lead Acetate Solution, Mercuric Chloride T.S., and Zinc Chloride Solution, but no precipitate with Platinic Chloride T.S. Ammonia Solution produces a white precipitate soluble in Chloroform or in an excess of the reagent, the latter producing a yellowish red solution. The aqueous solution yields a black precipitate of reduced metallic Silver when heated with Silver Nitrate Solution. Sulphuric Acid produces a yellow colour, turning slowly to green on warming, and finally to purple. Sulphuric Acid containing a trace of Selenious Acid gives a light bluish green coloration, gradually becoming dark green.

Not Official **GRINDELIA**

The Leaves and Flowering Tops of *Grindelia squarrosa*, Dunal, and *Grindelia robusta*, Nutt., from California.

It is now official in the *Ind* and *Col* Add for the Australasian and the North American Colonies.

Medicinal Properties—Antispasmodic, expectorant, slightly diuretic. Has been recommended in asthma, hay fever, bronchitis, whooping-cough, laryngismus stridulus, and cystitis.

Prescribing Notes—The Liquid Extract, whether made by U.S.P. or *Ind* and *Col* Add, has a peculiar, bitter, persistent taste, which requires a good deal of covering. The addition of Spirit of Chloroform, Syrup of Orange and Glycerin, is useful for this purpose. The so-called 'Alcoholic-Fluid Extract of Grindelia,' which is now introduced into the *Ind* and *Col* Add as *Extractum Grindelie Liquidum*, mixes more readily with Water, and makes a better looking and more palatable draught than either of the others.

Foreign Pharmacopœias—Official in *Fr* and U.S. Not in the others.

Descriptive Notes—The dried Leaves of *G. squarrosa*, Dunal, and *G. robusta*, Nuttall, are given. There is some difficulty in distinguishing these two species, since both have reflexed phyllaries. But as a rule the upper leaves of *G. squarrosa* taper towards the base, whilst those which are referred in the *P.B.* to *G. robusta* are broader towards the base and are somewhat shorter in proportion. The specific distinction depends on the nature of the achenes, which in *G. squarrosa* are 4-angled and without distinct auricular appendages, whilst those of *G. robusta* are bidentate. The species of the genus are very

variable, and it has been shown by Peniédés (*Ph. Jour* (4) 23, p 433) that the plant recognised in commerce as *G. robusta*, Nuttall, is really the *G. camporum* of Greene. It is this species that is now almost exclusively imported. When *Grindelia* was first introduced into this country it consisted of *G. squarrosa*. The Leaves of *G. squarrosa* are officially described in the *Ind* and *Col Add* as alternate, pale green, smooth, coriaceous, brittle oblanceolate or oblong lanceolate or elongate lanceolate, the lower leaves tapering considerably below. But these Leaves are no longer in commerce. The leaves of *G. robusta* are described as similar in texture and colour, but shorter and more oblong, with a cordate amplexicaul base, are furnished with ~~in~~ glandular hairs, and are sharply serrate at the margin. But these characters are those of the *Grindelia camporum* of Greene. In both species the reduced bracts in the involucre and the leaves are more or less covered with glossy patches of exuded resin. The taste is pungently aromatic and bitter, and the odour is balsamic.

Preparation

EXTRACTUM GRINDELIAE LIQUIDUM. LIQUID EXTRACT OF GRINDELIA

Percolate 20 of *Grindelia* with Alcohol (90 p c) until exhausted, distil off the Alcohol, and add to the residue 10 of Distilled Water and 2 of Sodium Bicarbonate, stir together, and after the Extract is dissolved and the effervescence is over, add Distilled Water to make 15, and finally Alcohol (90 p c), *q s* to yield 20 of product.

Dose —10 to 20 minims = 0.6 to 1.2 c c

This is official in the *Ind* and *Col Add* for the Australasian and the North American Colonies.

The official text directs the Sodium Bicarbonate to be previously added to the Distilled Water, but as it will not dissolve there is no point in it. This preparation deposits on keeping.

U. S. Fluid extract (1 in 1 w/v) by percolation with a mixture of Alcohol (95 p c) 3, Water, 1, Fr. Fluid Extract (1 in 1 w/w) with Alcohol (75 p c).

Not Official

EXTRACTUM GRINDELIAE —An Alcohol (90 p c) percolate, distilled and evaporated to an Extract. 100 of *Grindelia* yield 15 of Extract.

Dose —3 grains = 0.2 gramme, three times a day.

GUAIACI LIGNUM.

GUAIACUM WOOD

FR, GUAJACUM. GER, GUAJAKHOLZ, ITAL, LEGNO GUAJACO, SPAN, LENO DE GUAIACO.

The Heart-wood of *Guaiacum officinale*, L., or of *Guaiacum sanctum*, L.

It yields about 26 p c of Resin, consisting of Guaiacic, Guaiaconic and Guaiacinic Acid. It also contains two Saponins, a neutral Guaiac-saponin and Guaiacsaponic Acid.

Imported from St. Domingo and Jamaica.

Medicinal Properties.—See 'Guaiaci Resina'.

Foreign Pharmacopœias.—Official in all except Belg., Dan., Dutch, Fr., Hung. and Swed.

Descriptive Notes.—The official *Guaiacum Wood* consists of the dark-coloured heart-wood only, but in commerce it is usually met with in the form of turnings, containing more or less of the yellowish

sapwood, and sometimes of boxwood, or other woods used in turning. The turnings often require sifting to free them from powder. As Guaiacum Wood is heavier than Water and sinks in it, such admixtures can generally be separated by this means. The Wood is distinguished from other similar heavy dark greenish-brown wood by the medullary rays being one cell broad, and four, or sometimes three to six cells high (*P G*), by the usually solitary vessels with small pits, the numerous sphaeraphides of Calcium Oxalate, and the thick walled parenchyma with narrow lumen. It has a slightly acrid taste, and when heated a faintly aromatic odour. Although the use of the wood of *G. sanctum* is also permitted by the *P B*, no distinctive characters are given for it.

Test—Guaiacum Wood, when digested with Alcohol (90 p c) and filtered, yields a filtrate which gives, on the addition of diluted Ferric Chloride Test solution, a blue coloration. The ash varies from 1 to 2 p c.

GUAIACI RESINA

GUAIACUM RESIN

FR, RESINE DE GAIAC GER, GUAJAKHAR, ITAL, RESINA DE GUAIACO
SPAN, RESINA DE GUAIACO

The Resin obtained from the Stem of *Guaiacum officinale*, L, or of *Guaiacum sanctum*, L.

On dry distillation it yields Guaiacol similar to that found in Creosote.

Solubility—About 90 p c is soluble in Absolute Alcohol, Ether, Chloroform, Aromatic Spirit of Ammonia, and alkaline solutions, almost insoluble in Petroleum Spirit.

Medicinal Properties—Stimulant, diaphoretic, and alterative. It is employed in chronic forms of rheumatism and gout, especially in old people. It is used in acute tonsillitis, also in dysmenorrhoea, amenorrhoea, and syphilitic affections.

Generally prescribed in combination with other medicines.

It is innocuous, and might be taken for an indefinite period of time, and looked upon as a condiment rather than as a drug, as harmless as Ginger or any other condiment. Guaiacum possesses a considerable power, but less than Colchicum, in directly relieving patients suffering from gouty inflammation of any part, it might be given whenever there was but little fever. Guaiacum taken in the intervals of gouty attacks has a considerable power of averting their recurrence, in fact, it is a very powerful prophylactic. Guaiacum does not appear to lose its prophylactic power by long continued use—*L* '96, 1 1494, *B M J* '96, 1 1325.

In sub acute or chronic gout, in addition to Colchicum, 5 to 10 grains of the resin may very usefully be given in cachets two or three times daily. The cachets are far preferable to the tincture in a mixture, as the latter is nauseous and the precipitated resin tends to cling obstinately to the tongue and fauces—*A P Luff*, *Pr* '07, 1 166.

Confidence expressed in the efficacy of Guaiacum in many forms of chronic gout, in irregular gout, and also as a prophylactic of gout. It is best administered in the form of tablet, or as a cachet—*B M J* '00, 1 843.

10 grains in 1 tablespoonful of Malt Extract two or three times a day, before menstruation is expected, given to relieve the pain — *U. S. Disp.* 1105

Dose — 5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes — *Tragacanth* is better for the powder of *Guaiacum Resin* in Mixtures, *Mucilage of Acacia* is best for the Ammoniated Tincture — *Mucilage of Acacia*, 1 fl oz, Ammoniated Tincture, 6 fl dr, Water, to 6 fl oz

Incompatibles — Mineral Acid, Spirit of Nitrous Ether

Official Preparations — The Wood, used in the preparation of Liquor Sarsæ, of the Resin, *Mistura Guaiaci*, *Tinctura Guaiaci Ammonata*, *Trochiscus Guaiaci Resinæ*, used in the preparation of *Pilula Hydiargyri Subchloridi Composita*

Not Official — *Confectio Guaiaci Composita*, *Pulvis Guaiaci Composita*, *Tinctura Guaiaci*, and *Trochiscus Guaiaci*

Foreign Pharmacopœias — Official in Austr, Fr (Résine de Guaiac), Hung, Ital, Jap and Norw (Resina Guajaci), Mex, (Resina de Guayacan), Port, Span, Swed, Swis, and U S. Not in the others

Descriptive Notes — The Resin occurs in commerce in irregular masses, or in nearly globular tears varying in size from $\frac{1}{2}$ to 1 inch or more in diameter. The splinters of the Resin should be transparent, and of a yellowish-green or reddish-brown tint. The tear is the purest form; the Guaiacum in mass varying considerably in purity, the purest being obtained by cleaving the logs over a fire, and the inferior by boiling the chips in a solution of salt, some specimens of Guaiacum Resin in mass contain much woody matter and other impurities

Tests — Guaiacum Resin emits a balsamic odour when warmed, and possesses a slightly acid taste. A solution of the Resin in Alcohol (90 p c) yields on the addition of diluted Ferric Chloride Test-solution a blue coloration, and if the mixture be shaken with Chloroform the blue colour passes into the chloroformic layer. Paper moistened with the alcoholic solution becomes blue when exposed to the fumes of Nitric Acid. The percentage of matter insoluble in Alcohol (90 p c) should not amount to more than 10 p c. The impurities insoluble in Alcohol (90 p c) in good block Resin amount to 2.9 to 10 p c. The Acid value of crude lump Guaiacum varies from 90 to 95, the Alcohol-purified Resin from 90 to 100, the natural tears from 70 to 75. The *USP* gives the limits as not less than 70 nor more than 80. The *BP* gives neither the Acid value, the limit of matter insoluble in Alcohol (90 p c), nor the percentage of ash. The *USP* limit of matter insoluble in Alcohol (94.9 p c) is 15 p c and the ash not more than 4 p c. The ash of good commercial samples of the Resin varies from 1 to 4 p c., and should average 3.0 p c.

A standard of not less than 90 p c of matter soluble in Alcohol (90 p c) and not more than 3 p c of ash has been recommended.

The more generally occurring impurities are Colophony Resin, the similar but yellowish-brown Peruvian Guaiacum, and excess of woody fibre, Colophony may be detected by the very high Acid value. The *USP* macerates the powder with 4 or 5 times its weight of Petroleum Benzin, and requires that the filtrate should be colourless.

and that it should not give a green coloration on the addition of an equal volume of a 1 in 1000 Cupric Acetate Solution. Colophony and Peruvian Guaiacum may also be detected by dissolving the Resin in Chloroform and adding Bromine Solution. A blue coloration is yielded by the pure Resin, a red coloration by adulterated specimens. Excess of woody matter is indicated by the solubility in Alcohol (90 p c)

Preparations

MISTURA GUAIACI GUAIACUM MIXTURE

Guaiacum Resin, $\frac{1}{2}$ oz, Refined Sugar, $\frac{1}{2}$ oz, Tragacanth, in powder, 35 grains, mix these together intimately, then add gradually 20 fl oz of Cinnamon Water (1 in 40)

Tragacanth now used instead of Gum Acacia. As stated in previous editions of the *Companion*, not only does Tragacanth give a more diffusible mixture, but the colour does not change so rapidly, nor to the same extent as it does when Acacia is used.

Dose — $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

TINCTURA GUAIACI AMMONIATA AMMONIATED TINCTURE OF GUAIACUM

Add 4 oz of Guaiacum Resin in powder to $1\frac{1}{2}$ fl oz of Strong Solution of Ammonia, mixed with 16 fl oz of Alcohol (90 p c). After 48 hours, with occasional agitation, filter and add 30 minims of Oil of Nutmeg and 20 minims of Oil of Lemon. Wash the filter with Alcohol (90 p c) to make 20 fl oz of total product (1 in 5)

Dose — $\frac{1}{2}$ to 1 fl drm = 1 8 to 3 6 c c

Foreign Pharmacopœias — Official in U.S., 1 in 5 of Aromatic Spirit of Ammonia, Swed., Guaiacum Resin 3, Aqua Ammoniacæ (sp gr 0.960) 5, and Spirit 10, Port., Guaiacum Resin 3, Liquid Ammonia (sp gr 0.916) 3, Spirit 14, by weight. Not in the others.

Tests — Ammoniated Tincture of Guaiacum has a sp gr of 0.895 to 0.900, contains about 15 p c w/v of total solids and about 70 p c w/v of Absolute Alcohol.

TROCHISCUS GUAIACI RESINÆ GUAIACUM RESIN LOZENGE

3 grains of Guaiacum Resin in each, with Fruit Basis

Not Official

CONFECTIO GUAIACI COMPOSITA (*Syn* 'Chelsea Pensioner') —

Guaiacum, in powder, 1, Rhubarb, 2, Bitartrate of Potassium, 8, Sulphur, 16, one Nutmeg, Honey, 96 or q s — *Pharm. Form.*

Guaiacum Resin, 1, Rhubarb, in powder, 2, Acid Potassium Tartrate, $7\frac{1}{2}$, Nutmeg, in powder, 1, Sublimed Sulphur, $14\frac{1}{2}$, Clarified Honey, 74 — *B.P.C.*

Guaiacum Resin, in powder, $\frac{3}{4}$ oz, Mustard, 1 oz, Potassium Nitrate, in powder, $\frac{1}{2}$ oz, Rhubarb Root, in powder, $\frac{1}{2}$ oz, Sublimed Sulphur, 1 oz, Treacle, to 16 oz. Dose — 1 drm — *London.*

PULVIS GUAIACI COMPOSITUS ('Chelsea Pensioner') — Powdered Guaiacum Resin, Precipitated Sulphur, Heavy Magnesium Carbonate, Gum Acacia, Potassium Bicarbonate, of each equal parts. Dose — 20 to 40 grains — *St. George's.*

TINCTURA GUAIACI — Guaiacum Resin, 1, Alcohol (90 p c), 5

Dose — 30 to 60 minims = 1 8 to 3 6 c c

This has been incorporated in the *B.P.C.*

584 GUA [Solids by Weight, Liquids by Measure] Eng and US (Resin), 10 (Resin), all by weight

Foreign Pharmacopœias—Official in Austr, H, 1 in 5, Jap, Port and Swiss (Wood), 1 in 5, Fr, 1 in 5, for the presence of blood except US. Not in the others. Guaiacum Resin in each with

Along with Ozonic Ether it is employed as a test.
TROCHISCUS GUAIACI—2 grains of
Black Cumin Paste—*Throat*

Not Official
GUAIACOL—Obtained by fractional distillation of
Guaiacum Resin. It can also be obtained

A colourless, highly refractive liquid obtained from Wood Creosote. It has a characteristic aromatic odour (usually Beechwood) distilling from Guaiacum Resin. Synthetic Guaiacol is described as colourless rhomboid prisms, having an aromatic odour, which points to synthetic Guaiacol.

It forms the fraction of Wood Creosote between 200° and 205° C (392° and 401° F) 90, below 70° C. (1905) describes it as in colourless bottles of a dark amber tint and variety.

It should be preserved in well-closed glass bottles in all proportions with Alcohol (90 p c), Ether, Glycerin, and the fixed Oils (Almond and Olive).

Solubility—About 1 in 80 of Water, more used in place of Creosote in the (90 p c), Ether, Glycerin, and the fixed Oils (is better tolerated. Also given in

Medicinal Properties—Antiseptic, has also been used in erysipelas, internal treatment of phthisis, in which it is useful in sciatica, orchitis, and pleurisy. Olive Oil as an intralaryngeal injection. Exhaustion and profuse diaphoresis.

Disadvantages from continued use are: (1) in phthisis—B M J '96, 1 586
Applied externally: (2) toxic effects—L '98, 1 998

Hypocœmic: (under a Lanolin or Vaseline basis, in Large doses (60 minims) in phthisis with M J E '00, 1 92, '02, 1 20

A 10 to 20 p c Ointment made with L '99, 1 210

gonorrhœal epididymitis—T G '00, 145, 1 service in tuberculous cystitis—T G

Guaiacol vapour baths in bronchiectasis

Guaiacol (or the carbonate) of much use

'07, 313

Dose—1 to 5 minims=0.06 to 0.3 c. by given (mixed with Almond Oil) in

Prescribing Notes—It is generally used in mixtures with Glycerin and Water, and capsules, but it has also been given in M of Lavender, Oil of Cinnamon, or Compound Tincture of Gentian. But it can be treated in the same way as Creosote

both as regular Mixtures and Pills. in Belg, Dutch, Fr (Guaiacol), Ital, in the others

Foreign Pharmacopœias—Official 1116 to 1120. It boils at 205° C. Jap, Russ, Spain, Swiss and US. Not 205° C (892 and 401° F). Fr Codex

Tests—Guaiacol has a sp gr of 1.143, and the boiling point 205° C (401° F) and distils between 200° and 205° C. of Ferric Chloride T.S. added to a 1 in (1908) gives the sp gr at 15° C (59° F) a blue colour fading to green. It dissolves in twice its volume of 10 p c solution with 20 volumes of Water heating without residue. The sample may be shaken to a nearly white mass when the mixture should separate into two and Phenol. As a general test for the mixture should separate into two with twice its volume of Petroleum Ether. The mixture should separate into two clear layers, any turbidity or failure to separate may be taken as an indication of the presence of impurities. The failure of the solid mass to dissolve in Hydroxide solution test described above, also indicates the presence of

produce a clear solution with 20 times its volume of Water indicates the presence of oily hydrocarbons Creosote may be detected by the reddish colour produced when the specimen is treated with 10 times its volume of Sulphuric Acid Guaiacol develops a pure yellowish colour It should leave no residue on volatilisation

NEBULA GUAIACOL ET MENTHOL —Guaiacol, 10 minims, Menthol, 60 grains, Paraffin Liquid, to 1 fl oz —*A Ph F*

Guaiacol, 2, Menthol, 4, Liquid Paraffin, *q s* to produce 100 —*B P C*

VASOLIMENTUM GUAIACOLI —Guaiacol, 20, Liquid Vasoliment, 80 —*Hager*

Parogen Guaiacolis *Syn* Guaiacol Vasoliment —Guaiacol, 20, Parogen, 80 —*B P C*

GUAIACOL (Synthetic) C_9H_8O , eq 123.13 —A crystalline substance which melts at about $28^{\circ}C$ ($82.4^{\circ}F$), but frequently remains liquid much below this temperature It is said to yield more uniform results than the ordinary medicinal liquid Guaiacol, which is not so definite in composition Soluble 1 in 50 of Water

It should be kept in well closed glass bottles of a dark amber tint and protected as far as possible from the light

Dose —1 to 5 grains = 0.06 to 0.32 gramme

Tests —Synthetic Guaiacol melts at $28^{\circ}C$ ($82.4^{\circ}F$), and when melted should answer the tests and be free from the impurities mentioned under Guaiacol

GUAIACOL BENZOATE Benzosol $C_9H_7O \cdot C_6H_5O$, eq 226.38 —A white crystalline powder, having an aromatic taste and odour It contains theoretically 54.39 p c of Guaiacol Almost insoluble in Water A non irritating form of Guaiacol, recommended in phthisis and in diabetes —*MP '94*, 1 269 *L* 96, 11 551

Dose —5 to 10 grains = 0.32 to 0.65 gramme, usually given in cachets or tablets

Tests —Guaiacol Benzoate melts at about $56^{\circ}C$ ($132.8^{\circ}F$) and when prepared from synthetic Guaiacol at $59^{\circ}C$ ($138.2^{\circ}F$) It is decomposed by Alcoholic Potassium Hydroxide Solution (Semi normal) and may be volumetrically determined by means of this solution A weighed quantity of 1 gramme is dissolved in 25 c c of Alcohol (90 p c) mixed with 25 c c of Semi normal Volumetric Alcoholic Potassium Hydroxide Solution, and saponified under a reflux condenser The excess of Semi normal Alkali Solution is titrated with Semi normal Volumetric Hydrochloric Acid Solution, and the amount of Semi normal Volumetric Alkali Solution absorbed is calculated into Guaiacol Benzoate, 1 c c of Semi normal Volumetric Potassium Hydroxide Solution is equivalent to 0.11819 gramme of the pure salt A solution in Alcohol (90 p c) should yield no appreciable coloration with Ferric Chloride T.S. It leaves no weighable residue when ignited with free access of air

GUAIACOL CAMPHORATE (Guacamphol) —Colourless needles or a white or nearly white powder, having an aromatic odour Insoluble in Water, soluble in cold, readily soluble in hot Alcohol (90 p c), and in Chloroform

Used with success in the night sweats of phthisis —*C D '01*, 11 344

Dose —5 grains = 0.32 gramme

GUAIACOL CARBONATE Duotal $(C_9H_7O)_2CO_3$, eq 272.05 —A white crystalline powder, carbonaceous and tasteless It contains theoretically 90.5 p c of Guaiacol

Solubility —Insoluble in Water, about 1 in 70 of Alcohol (90 p c)

A non irritating form of Guaiacol in phthisis —*B M J E '92*, 1 8, '93, 11 83, '95, 1 8, *L* '96, 11 1374, '98 1 222, 960

Dose —3 to 10 grains = 0.2 to 0.65 gramme, which may be gradually increased to 60 grains = 4 grammes

Rheumatoid arthritis, whether acute or chronic, is of infective origin, and

infection is believed to take place from the alimentary tract. Intestinal antiseptics, *e g*, Guaiacol Carbonate, are stated to possess a high value. The great value of the drug is corroborated, but not attributed to its antiseptic action in the intestine — *L* '05, 1 718

It has the advantage of being less disagreeable than Creosote, and practically tasteless, but is much more expensive — *Edin Med Jour* '05, 463

The most convenient form of administering Guaiacol is the Carbonate in cachets. In rheumatoid arthritis, at first from 5 to 10 grains should be given three times a day, and the dose should be increased by 1 to 2 grains each week until from 15 to 20 grains are being taken in each dose. It is essential that this treatment should be continued for at least twelve months. The beneficial effects of the Guaiacol are very much increased by administering at the same time a mixture containing Potassium Iodide, the depressing effect of the Iodide should be counteracted by its combination with tonics — *A P Luff, B M J* '07, 11143

Foreign Pharmacopœias — Official in Austr, Belg, Fr, Ital, Jap, Russ, Swiss and U S

Tests — Guaiacol Carbonate melts at about 84° C (183 2° F). When heated with Alcoholic Potassium Hydroxide Solution (about 3 p c w/v) yielding Guaiacol when the liquid is acidified. No bluish-green is produced on the addition of one or two drops of Ferric Chloride T S to its solution in Alcohol (90 p c). It should leave no weighable residue on ignition.

GUAIACOL CINNAMATE Styraol $C_{15}H_{12}O$ $C_9H_7O_2$, eq 252.20 — Colourless, crystalline needles, almost insoluble in Water, soluble in Alcohol (90 p c) and in Chloroform. It contains theoretically 48.8 p c of Guaiacol. Recommended in phthisis, and also in cystitis and gonorrhœa.

It is tasteless, and does not split up into its constituents until it has passed through the pylorus. Very useful where intestinal tubercle is suspected, or where there is troublesome diarrhœa. Most serviceable in large cavities, with offensive sputum and fetid breath. Appears to be more beneficial than Guaiacol. Employed in form of powder or tablets, the latter to be bitten into minute particles lest they pass through the intestine unchanged — *F T* '07, 90

Dose — 5 grains = 0.32 gramme, 3 times daily

Tests — Guaiacol Cinnamate melts at 130° C (266° F). It should yield no weighable residue when heated with free access of air.

GUAIACOL PHOSPHATE — A white crystalline powder, insoluble in Water, soluble in Alcohol (90 p c) and in Chloroform. Useful in tuberculosis and in typhoid fever — *L* '02, 1 1711

Dose — 1½ to 3 grains = 0.1 to 0.2 gramme three or four times daily

There is also a crystalline Guaiacol Phosphite, dose, 5 to 10 grains = 0.32 to 0.65 gramme

GUAIACOL VALERIANATE (Geosote) — A yellowish, oily liquid, almost insoluble in Water. Used in tuberculous, bronchial affections and diarrhœa — *L* '97, 11 932, *B M J E* '98, 1 75, *P J* '97, 1 425

Dose — 2 to 3 minims = 0.12 to 0.18 c c or more

Tests — Guaiacol Valerianate has a sp gr of about 1.037. It boils at 245° to 265° C (474° to 509° F). It should leave no weighable residue when ignited with free access of air.

GUAIACETIN (Sodium Pyrocatechin-monoacetate) — A white crystalline powder, having a faint odour and taste of Guaiacol. Soluble in Water, insoluble in Alcohol (90 p c). Recommended in tuberculosis — *Pr* lxix 704

Dose — 4 to 8 grains = 0.25 to 0.5 gramme three or four times daily.

GUAIACYL (Calcium Oxyguaiacol-sulphide) — A greenish or greyish-mauve powder. Readily soluble in Water and in Alcohol (90 p c). A 5 to 10 p c solution is useful as a local anæsthetic.

Dose — 0.5 to 1.5 c c of a 5 p c solution, 1 c c of a 10 p c solution

GUAIIFORM (Geoform) — A yellow or brownish-yellow, tasteless powder, insoluble in Water, soluble in Alcohol (90 p c), and in Ether. Stated to be a

non-irritating preparation, and likely to be of use in pulmonary tuberculosis and typhoid fever. The Tannic Acid compound is known as 'Tannoguaiacform'—*L* '02, i 912, *P J* '02, 61

THIOCOL (Potassium Guaiacol Sulphonate)—White, glistening crystals. Readily soluble in Water, insoluble in Alcohol (90 p c). Recommended in phthisis, stated not to irritate—*L* '99, i 240, *B M J E* '01, i 16, *P J* '01, ii 645

Dose—10 to 20 grains = 0.65 to 1.3 grammes three times a day

The somewhat bitter taste of Thiocol may be disguised by Syrup of Orange. A Syrup containing 5 grammes of Thiocol in each 100 grammes is known under the name of 'Sirolin'.

Aphthisin is stated to be a mixture of Potassium Guaiacol Sulphonate and Ammonium Sulphichthyolate—*P J* '02, ii 137

Among the various other compounds containing Guaiacol which have received attention in medical literature are **Euguforn** (Acetyl methylenediguaiacol), a greenish white powder, insoluble in Water, antiseptic and anaesthetic, recommended as a dusting powder, also a 50 p c solution in Acetone. **Guaiacol Cacodylate**, a dangerously unstable salt recommended subcutaneously in $\frac{1}{4}$ to $\frac{1}{2}$ grain doses in tuberculosis. **Guaiakinol** (Quinine di bromo guaiacolate), yellow crystals, readily soluble in Water, recommended for external use in erysipelas. **Quaiquin** (Quinine Guaiacol bi sulphonate), a yellow powder, readily soluble in Water, introduced as a substitute for Guaiacol. **Guaiamar** (Glycerol ester of Guaiacol), a white, non hygroscopic crystalline powder, used as an antiseptic (dose, 5 to 10 grains). **Guaiasanol** (Diethylglycocol Guaiacol), a white crystalline powder, readily soluble in Water, used as an antiseptic. **Guaiacol Salol** (Guaiacol Salicylate), a white crystalline powder, insoluble in Water, soluble in Alcohol (90 p c), recommended in phthisis

Not Official GUARANA

The Seeds of *Paullinia Cupana*, H B and K, dried in the sun, and then roasted and reduced to a fine powder, this is moistened with a little Water, exposed to the night dew, and when it has become a hard paste is rolled into cylinders, these are further dried in the sun or in the chimneys of the huts. It is exported from Brazil.

True Guarana is very hard, heavy, and, when powdered, is reddish grey, whilst the sophisticated is much lighter in colour, it contains about 4 p c of an alkaloid **Guaranine** (dose, 1 to 5 grains = 0.06 to 0.32 grammes), generally considered to be identical with Caffeine.

The *U S P* requires that it shall yield, when assayed by the process outlined below, not less than 3.5 p c of its alkaloidal principles.

Medicinal Properties—Nervine tonic. It is used chiefly for curing sick headache, but is also useful in diarrhoea, dysentery, and as a tonic and stomachic in convalescence.

Dose—10 to 60 grains = 0.65 to 4 grammes infused in boiling Water and sweetened, and repeated if necessary in two hours.

Foreign Pharmacopœias—Official in Austr., Hung., Ital., Mex., Port., Span., Swiss and U S.

Tests—Guarana is required by the *U S P* to yield a definite percentage of alkaloidal principles. The following is an outline of the *U S P* method of determination.—A weighed quantity of 6 grammes of the specimen in No. 60 powder is shaken in an Erlenmeyer flask, at intervals, for half an hour, with 120 c c of Chloroform and 6 c c of Ammonia Solution. The mixture is allowed to stand for four hours, and is then filtered, a measured quantity of 100 c c (= 5 grammes Guarana) is collected, and the Chloroform distilled off in a water-bath. The residue is dissolved in a mixture of 2 c c of Normal Volumetric Sulphuric Acid Solution and 20 c c of warm Water. The cooled liquid is filtered into a separator, the flask and filter are washed with Water and the washings transferred

to the separator, 2 c c of Ammonia Solution added, and the alkaloids extracted by shaking the solution with 20 c c of Chloroform, the extraction being repeated with two separate portions each of 10 c c of Chloroform. The separated chloroformic solutions are mixed, the Chloroform distilled, 2 c c of Ether is added to the dry residue, the Ether carefully evaporated on a water-bath, and the residue dried at this temperature till constant in weight. The weight of residue multiplied by 20 gives the percentage w/v of alkaloids.

Preparations

ELIXIR GUARANÆ—Guarana, in No. 60 powder, 4 oz., Light Magnesia, $\frac{1}{2}$ oz., Oil of Cinnamon, 6 minims, Syrup, 2 fl. oz., Alcohol (60 p c), q s to produce 20 fl. oz.—*B P C Formulary* 1901 incorporated in the *B P C*.

Dose—30 to 120 minims = 1.8 to 7.1 c c.

The *B P C Supplement* has altered the Light Magnesia to 'Purified Talc or Kaolin'.

FLUIDEXTRACTUM GUARANÆ (U S)—Guarana, in No. 60 powder, percolated with Alcohol (49 p c), and treated in the usual manner to make 100 c c of Fluidextract.

Average Dose—30 minims (about 2 c c).

Fluidextractum Guaraniæ *U S P* is required to contain 3.5 grammes of the alkaloids from Guarana in 100 c c.

This has been incorporated in the *B P C Supplement*, using Alcohol (45 p c).

Tests—The *U S P* method of determining the alkaloids in this Fluidextractum may be briefly outlined as follows.—A measured quantity of 5 c c of the Fluid Extract is well shaken in a separator with 15 c c of Chloroform and 1 c c of Ammonia Solution, the shaking being repeated with two separate portions each of 10 c c of Chloroform. The chloroformic liquids are separated, mixed, and evaporated to dryness. The residue is dissolved in a mixture of 2 c c of Normal Volumetric Sulphuric Acid Solution and 20 c c of warm Water. The cooled solution is transferred to a separator, the vessel and filter washed with Water, and the alkaloids are extracted from the mixed solution and washing by shaking with 20 c c of Chloroform and 2 c c of Ammonia Water. The extraction is repeated with two separate portions each of 10 c c of Chloroform. The separated chloroformic liquids are mixed, the Chloroform removed by evaporation, the dry residue mixed with 2 c c of Ether, and this in turn is carefully removed by evaporation. The residue is dried till constant in weight at the water-bath temperature, and weighed when cool. This weight, multiplied by 20, yields the percentage w/v of alkaloids in the Fluid Extract.

TINCTURA GUARANÆ—Guarana, in fine powder, 1, Alcohol (60 p c), q s. to produce 4.

Dose—30 to 120 minims = 1.8 to 7.1 c c.

This has been incorporated in the *B P C*, employing Alcohol (90 p c), but in the *B P C Supplement* this has been altered to Alcohol (60 p c).

Not Official

GUMMI INDICUM.

INDIAN GUM

A gummy exudation from the Wood of *Anogeissus latifolia*, Wall., is official in *Ind* and *Col*. Add for India and the Eastern Colonies, and is used in official preparations for which Gum Acacia is used, the former being taken for every two parts ordered of the latter.

GUMMI RUBRUM.—See EUCALYPTI GUMMI

Not Official

GUTTA PERCHA

Tough, somewhat flexible pieces, of a light brown or chocolate colour, which become hard and brittle on keeping, but can be softened again in warm Water

The concrete Juice of *Dichopsis Gutta*, and of several other trees of the natural order Sapotaceæ

It was official in *B P* '85, but is replaced in *B P* '98 by Caoutchouc, a solution of which is now used for Charta Sinapis

Solubility—Almost entirely soluble in Chloroform, yielding a more or less turbid solution Entirely soluble in Oil of Turpentine, Carbon Bisulphide, and Benzol Insoluble in Water, Alcohol, alkaline solutions, or dilute acids

Medicinal Properties—Used for making splints as Gutta Percha tissue for keeping surgical dressings moist, as a solution for mixing with medicaments for chronic skin diseases, and applying like Collodion

Foreign Pharmacopœias—Official in Belg, Fr, Ger (also Percha Lamellata), Hung, Jap (also Gutta Percha Depurata), Port, Span, Swed (also Gutta Percha Laminata), and Swiss, which has also Percha Lamellata

LIQUOR GUTTA PERCHA—Gutta Percha, in thin slices, 1, Chloroform, 8, Lead Carbonate, in fine powder, 1 Add the Gutta Percha to 6 of the Chloroform in a stoppered bottle, and shake them together frequently until solution has been effected Then add the Lead Carbonate previously mixed with the remainder of the Chloroform, and having several times shaken the whole together, set the mixture aside, and let it remain at rest until the insoluble matter has subsided Lastly, decant the clear liquid, and keep it in a well stoppered bottle—*B P* '85

Traumaticine—A solution of 1 of Gutta Percha tissue in 10 (by weight) of Chloroform It produces a thin delicate film when painted on the skin, and causes neither tension nor pain It is used for medicated applications—*P J* (3) xiv 341 A vehicle for the administration of Mercury in syphilis—*L* '94, ii 590

B P C uses 1 of Gutta Percha in 10 of Chloroform by weight, the same as **Traumaticine**, and the directions for making the solution are those of *B P* '85

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Mex, Span and Swiss, Gutta Percha 1, Chloroform 9 (by weight), all have Traumaticine either as a title or as a synonym Jap (Liquor Guttaperchæ) 1 and 10, with Lead Carbonate

UNNA'S PLASTER MULLS consist of a very thin sheet of Gutta Percha coated on one side with an adhesive substance (Aluminium Oleicum) containing one or more medicinal substances, and backed on the other side with Mull (undressed muslin)—*L* '86, ii 575

Not Official

GYNOCARDIÆ OLEUM

Prior to 1900 it was supposed that the Chaulmoogra Oil of commerce was obtained from the seeds of *Gynocardium odorata*, but it was pointed out by Holmes, on the authority of Dr Plam, that Chaulmoogra Seeds and Oil are the produce of *Taraktogenos Kurzii*, King Power and Barrowcliff have extracted and examined the Oil from seeds of *Gynocardia odorata* supplied to them by Mr David Hooper

Gynocardia Oil consists, according to the above-named authors, of the glyceryl esters of the following Acids (1) Linolic or isomerides of the same series, (2) Palmitic Acid in considerable amount, (3) Linolemic and Isohnolemic Acids, the latter preponderating, and (4) Oleic Acid in relatively small amount The seeds also contain 5 p.c. of a crystalline glucoside, Gynocardin ($C_{11}H_{19}O_5N$, $1\frac{1}{2}H_2O$, eq 357.51) and a hydrolytic Enzyme, Gynocardase

Tests—Gynocardia Oil has, according to Power and Barrowcliff, a sp gr of 0.925 at 25° C (77° F). It is optically inactive. It has an Acid value of 4.9, a Saponification value of 197.0, and an Iodine value of 152.8. The oil extracted from the seeds by Ether has a sp gr of 0.927 at 25° C (77° F), an Acid value of 5, a Saponification value of 199.6, and an Iodine value of 152.

CHAULMOOGRA OIL—Chaulmoogra Oil of commerce is obtained from the Seeds of *Taraktogenos Kurzii*, King, a plant which is a native of Burma. The shells, which were separated from the fresh Chaulmoogra seeds by Power and Gornall, represented 34 p.c. of their weight, the kernels yielded, by expression, an amount of fixed oil corresponding with 30.9 p.c. of the entire seeds. A portion of the kernels, when completely extracted with Ether, yielded 55 p.c. of their weight of fixed oil, corresponding with 38.1 p.c. of the entire seed (having 30.7 p.c. of shells). It is a soft solid, having a faintly yellow colour and a characteristic odour.

The Oil prepared by these authorities from the Seeds yielded, on hydrolysis, a substance having the formula and m.p. of Phytosterol, Glycerol, and a mixture of fatty acids having a m.p. of 44° to 45° C (111.2° to 113° F), an optical rotation in Chloroform Solution of +52.6°, an Acid value of 215, and an Iodine value of 103.2. Palmitic and Chaulmoogric Acid were identified in this mixture.

The Oil has been long known and used in India, it has a disagreeable taste and smell, and can be readily melted by a gentle heat.

Oleum Gynocardiæ is official in the *Ind* and *Col Add* for India and the Eastern Colonies, with the synonym Chaulmoogra Oil.

Medicinal Properties—Recommended in leprosy, also as an external application in psoriasis, obstinate eczema, and other skin diseases, chronic rheumatism, and gout, and in phthisis.

In leprosy—*B.M.J.E.* '93, ii, 4, '01, ii, 79, *L.* '07, ii, 1515.

4 minims in capsule three times daily in leprosy, dose increased until 50 capsules per diem were taken—*L.* '02, ii, 1196.

Dose—5 to 10 minims = 0.3 to 0.6 c.c., gradually increased to 30 to 60 minims = 1.8 to 3.6 c.c. three or four times a day, should be given after meals in Milk or emulsion with Gum Acacia, or better still in capsules.

Tests—The Oil has, according to the above-named authors, a sp gr of 0.951 at 25° C (77° F), or of 0.940 at 45° C (113° F). It is dextrogyrate, the optical rotation being +52° in a tube of 100 mm. The m.p. is 22° to 23° C (71.6° to 73.4° F). It has an Acid value of 23.9, a Saponification value of 213.0, and an Iodine value of 103.2.

The Ether-extracted oil has a sp gr of 0.952 at 25° C (77° F), or of 0.942 at 45° C (113° F). It is dextrogyrate, the optical rotation being +51.8° in a tube of 100 mm. The m.p. is 22° to 23° C (71.6° to 73.4° F). It has an Acid value of 20.8, a Saponification value of 208, and an Iodine value of 104.4.

A specimen of Chaulmoogra Oil, which had been in stock for some considerable time, examined in the author's laboratory, gave an Acid value of 29.4, an Iodine value of 163, a Saponification value of 197.4, and an Iodine value of 99.06, the oil yielded 99.56 p.c. of fatty acids, having a combining weight of 288.

Chaulmoogric Acid ($C_{18}H_{32}O_2$, eq. 214.38), isolated by Power and Gornall from the Chaulmoogra Oil described above, has a m.p. of 55° C (131° F), is dextrogyrate, the optical rotation being +56°. It is soluble in Ether, Chloroform, Carbon Tetrachloride, or by Nitric Acid. It is also soluble in concentrated Sulphuric Acid, with much decomposition, forming Sulphur Dioxide.

Magnesium Gynocardate—A granular powder.

Dose—1 to 3 grains = 0.06 to 0.2 gramme.

UNGUENTUM GYNOCARDIÆ (*Ind* and *Col Add*)—A 10 p.c. Ointment of Gynocardia Oil in a mixture of 4 of Hard and 5 of Soft Paraffin.

For India and the Eastern Colonies.

HÆMATOXYLI LIGNUM.

LOGWOOD

FR, BOIS DE CAMPECHE, GER, BLAUHOLZ, ITAL, CAMPEGGIO, SPAN,
PALO DE CAMPECHEThe Heart-wood of the trunk of *Hæmatoxylon Campechianum*, L

Imported from Campeachy in Central America, from Honduras and Jamaica, that from Campeachy being the most valuable

Medicinal Properties —Astringent, without irritating properties, useful in diarrhoea of phthisis and chronic diarrhoea and dysentery, and in passive hæmorrhages, in infantile diarrhoea, it does not tend to cause subsequent constipation Also as an injection for leucorrhœa It colours the urine and fæces dark red**Incompatibles** —Mineral Acids, metallic salts, Lime Water, Tassar Emetic**Official Preparation** —Decoctum Hæmatoxyl**Not Official** —Extractum Hæmatoxyl, Extractum Hæmatoxyl Liquidum, Hæmatoxylum, and Hæmatein**Foreign Pharmacopœias** —Official in Austr, Mex (Palo de Campeche), Port (Campeche), U S Not in the others**Descriptive Notes** —Logwood consists of the heart-wood of the trunk of *Hæmatoxylon Campechianum*, a leguminous tree indigenous in Central America There are several varieties of the tree, four being recognised in Honduras and three in Jamaica, the wood of which varies in tinctorial power The kinds imported from Campeachy and San Domingo are considered the best The heart-wood of the tree only is used, the bark and sapwood being removed It is imported in logs about 3 feet long, externally often dark purplish red, and reddish or orange-brown internally In retail commerce it is sold in chips or, more rarely, in coarse powder, and for dyeing purposes is usually fermented from four to six weeks by moistening it and exposing it to the air During this process the Hæmatoxylum, which in the pure state is colourless, becomes oxidised in the presence of atmospheric Ammonia to Hæmatein, the presence of which is recognised by the bronzy-green midescence observable on the surface of the chips The unfermented wood is official for use in medicine, and is described as being purplish-red externally, and internally reddish-brown with medullary rays 4 cells wide (*U S P*) When chewed it colours the saliva pink It should have a slight, agreeable odour and a sweetish, astringent taste The odour recalls that of violets, and is perceptible in the decoction The wood contains about 9 to 12 p c of Hæmatoxylum An extract of Logwood is prepared for technical purposes which resembles Kino in appearance, but is easily distinguished by its sweet taste The only wood with which it is likely to be confounded is Brazil wood, which gives a red, not blue, colour with alkalis, and gives Picric Acid when boiled with Nitric Acid, whilst Logwood gives only Oxalic AcidSee also *P J* (4) vi 284**Tests** —Hæmatoxylum Wood when ignited with free access of air should not leave more than 2 p c of ash

Preparation

DECOCTUM HÆMATOXYLI DECOCTION OF LOGWOOD

Boil 1 oz of Logwood, in chips, with 24 fl oz of Distilled Water, adding 70 grains of bruised Cinnamon Bark towards the end of the process, strain, and wash with Distilled Water to make 20 fl oz

(1 in 20)

Iron vessels should not be used

Dose.— $\frac{1}{2}$ to 2 fl oz = 14 2 to 56 8 c c

Not Official

EXTRACTUM HÆMATOXYLI (B P 1885) —Logwood, in fine chips, 1, boiling Distilled Water, 10, infuse 24 hours, boil to 5 strain and evaporate to dryness by a water bath, stirring with a wooden spatula Iron vessels should not be used

Dose —10 to 30 grains = 0 65 to 1 94 gramme

This has been incorporated in the B P C

Foreign Pharmacopœias —Official in U S Not in the others

EXTRACTUM HÆMATOXYLI LIQUIDUM —Boil 20 of *Unfermented* Logwood, in No 16 powder with 40 of Distilled Water for half an hour, and strain, repeat the process with 40 more of Water, and again for the third time, and having mixed the strained liquors, evaporate over a water-bath (or in *vacuo*) to the measure of 17 and add 3 of Alcohol (90 p c), allow it to settle for a week, then draw off the clear liquor from the sediment

Dose —30 to 120 minims = 1 8 to 7 1 c c

The above B P C *Formulary* 1901 general process has been incorporated in the B P C, except that Logwood is in No 20 powder instead of No 16, and the product is made up to a volume of 1 in 1

HÆMATOXYLIN $C_{16}H_{11}O_8$, eq 299·84 —Bright yellow prismatic or granular crystals, sometimes brownish externally It possesses a sweet taste somewhat resembling Liquorice The prismatic crystals contain 3 molecules of Water of crystallisation, the granular crystals 1 molecule Sparingly soluble in cold Water, readily in Alcohol and Ether It is also soluble in solutions of the fixed and volatile alkalis with the production of solutions which rapidly acquire a purple colour It has the characters of a weak acid, and unites with basic ions to form compounds, which are colourless when perfectly pure, but soon pass into strongly coloured products, owing to the great avidity with which they absorb atmospheric Oxygen Used as a nuclear stain for histological and pathological sections

Tests —Hæmatoxylin loses part of its Water of crystallisation at 100° C (212 F) but the remainder only at 110° to 120° C (230° to 248° F) It fuses upon further heating, about 110° to 120° C (230° to 248° F) till higher temperature decomposes, leaving a bulky carbonaceous residue A solution yields with neutral or basic Lead Acetate a bluish-white precipitate rapidly darkening when exposed to the atmosphere Chloride Solution a permanent rose-coloured precipitate Solution of Alum yields a bright red colour, but no immediate precipitate A dilute Acetate Solution yields a fine purple precipitate Ferric Chloride Acid and Potassium Bichromate, and its solution reacts with Potassium Tartrate (Fehling's) Solution and Silver Nitrate Solution

Foreign Pharmacopœias —Official in Belg and Jap

HÆMATOXYLIN SOLUTION See Indicators of Neutrality

HÆMATEIN $C_{16}H_{12}O_8$, eq 297·84 —A brownish-red powder sparingly soluble in Water the atmospheric oxidation of an ammoniacal solution of its ammonium salt of Hæmatin being decomposed by

Acetic Acid The Ammonium salt forms a deep violet crystalline powder exhibiting a metallic lustre. It is soluble in Water, the solution readily reducing Silver Nitrate Solution. It yields with Copper Sulphate Solution a violet blue precipitate, and with Stannous Chloride Solution a violet precipitate.

Not Official

HÆMOGLOBIN

The substance to which in one or other of its modifications the blood owes its colour, and the chief solid constituent of the red blood corpuscles. Has been given with considerable success in the treatment of anæmia. It readily combines with free Oxygen to form oxyhæmoglobin or hæmato crystallin. It has been prepared in the form of crystals, but its preparation in this form is attended with some difficulty on account of its ready solubility in Water. A colloidal form is also known as colloidal hæmoglobin.—*L* '02, i 910, *B M J* '02, i 738

It occurs in commerce as an **Extract** (Pfeuffer's), in **Scales** (Merck) and as a dry powder, **Sanguis Bovinus Exsiccatus**, defibrinated and desiccated ox blood.

HÆMATOGEN—An aromatic fluid preparation, stated to contain pure hæmoglobin, the salts of the blood, the albuminous constituents of the serum, and glycerin.—*L* '99, ii 388

Under the name of **Sicc**, a solid preparation of hæmatogen has been introduced. It is a brownish black powder, soluble in Water.

LIQUOR HÆMOGLOBIN CO (Vinsip)—A fluid preparation, stated to contain hæmoglobin, and the albuminous constituents of the blood.—*L* '01, ii 735

HÆMOL—A dark brown powder, slightly soluble in Water, produced by the action of reducing substances, e.g., Zinc dust, on the colouring matter of the blood.

Dose—3 to 8 grains = 0.2 to 0.52 gramme

Under the name of **Ferrohæmol**, **Cuprohæmol** and **Zincohæmol**, compounds containing respectively Iron, Copper and Zinc with Hæmol have been introduced, **Bromo-hæmol** has been used in the treatment of epilepsy.

HÆMOGALLOL—A dark brown or reddish brown amorphous powder, slightly soluble in Water. Produced by the action of Pyrogallol on the colouring matter of the blood.

Dose—1 to 5 grains = 0.06 to 0.32 gramme

HAMAMELIS.

HAMAMELIS

Both the dried Bark, and the fresh and dried Leaves of *Hamamelis Virginiana*, L., are official.

Medicinal Properties—A local astringent and hæmostatic. Used in epistaxis, hæmatemesis, bleeding piles, and other conditions in which tannin is used.

Prescribing Notes—For local application, 1 of the Tincture is diluted with 10 or 20 of Water or the Liquor with 1 or 2 of Water. The ointment is used for piles, as is also a suppository of Hamamelin.

When equal Volumes of Tincture of Hamamelis and Tincture of Hydrastis are mixed, a precipitation will occur unless each Tincture be mixed with an equal Volume of Glycerin.

Official Preparations—Of the Bark, Tinctura Hamamelidis, of the Dried Leaves, Extractum Hamamelidis Liquidum, of the Fresh Leaves, Liquor Hamamelidis, of the Liquid Extract, Unguentum Hamamelidis

Not Official—Extractum Hamamelidis, Hamamelidis, Pasta Hamamelidis, Suppositorium Hamamelidis, Hamamelidis, and Hamamelin

Descriptive Notes—Hamamelis leaves are official in the *BP*, both fresh and dried, but in the *USP* only the dried leaves, collected in autumn. The dried leaves are more or less broken in commerce, but the fresh leaves are broadly oval, 3 to 6 inches (7 to 15 cm long) (10 cm *USP*), shortly stalked, cordate and unequal at the base, and sinuate at the margin, pinnately veined, paler below, with prominent veins furnished with stellate hairs, and an astringent taste, with slight bitterness. It has been found that the leaves contain more tannin in the autumn, and that the cells of the hairs have thicker walls, a dark line often marking the lining of the cell in the autumn, the walls becoming yellow, and the granular and oily contents. The odour of the distillate of the leaves is quite characteristic and is apparently the result of decomposition of the volatile oil, and is not perceptible in the dried leaves.

HAMAMELIDIS CORTEX. HAMAMELIS BARK *BP Syn* — WITCH HAZEL BARK

The dried Bark of *Hamamelis Virginiana*

Foreign Pharmacopœias—Official in Mex, Span and *US*

Descriptive Notes—Hamamelis Bark occurs in commerce in thin quilled pieces of pale brownish-buff or fawn colour, the outer surface or cork being thin, of a greyish tint, cracking and forming scales, and easily exfoliating, so that the inner bark, which is cinnamon-coloured or reddish-brown, often occurs free from it in commerce. The transverse fracture is short externally, but laminated internally with weak fibres. The taste is faintly astringent and somewhat mucilaginous. Its activity is apparently due chiefly to a volatile oil, as it only contains 8 to 10 p.c. of Tannin and a small quantity of bitter principle. Hamamelis Bark is about $\frac{1}{8}$ inch (1.5 mm) thick, *BP* (1 to 2 mm *USP*), 2 to 8 inches long (0.5 to 2 dm). The transverse section exhibits a complete ring of sclerenchymatous cells near the outer surface and numerous tangentially elongated bundles of bast fibres. Willow Bark bears some resemblance to Hamamelis Bark. It has a dull greyish-brown cork, is usually striated or wrinkled on the outer surface and does not exhibit a line of sclerenchymatous cells, and the bast fibres are much tougher than those in Hamamelis Bark, the taste also is more astringent.

Tests.—The Bark yields about 5 p.c. of ash and the amount yielded should not be much in excess of this figure.

An ash limit is stated not to be necessary for inclusion in the *BP*.

Preparations.

TINCTURA HAMAMELIDIS. TINCTURE OF HAMAMELIS

2 of Hamamelis Bark, percolated with Alcohol (45 p.c.) to yield 20 (1 in 10)

Dose —30 to 60 minims = 1 8 to 3 6 c c

Foreign Pharmacopœias —Official in Fr, 1 in 5 from leaves prepared with Alcohol (60 p c), Mex, 1 in 5, and Span, Bark 1 and Leaves, 1 in 20

Tests —Tincture of Hamamelis has a sp gr of 0 950 to 0 955, contains about 2 0 p c w/v of total solids and about 49 0 p c v/v of Absolute Alcohol

Not Official

EXTRACTUM HAMAMELIDIS —Hamamelis Bark in powder, percolated with Alcohol (60 p c) and the percolate evaporated to the consistence of an extract
Yield of Extract, 20 to 25 p c

Dose — $\frac{1}{2}$ to 2 grains = 0 032 to 0 13 gramme in pill

$\frac{1}{2}$ grains = 0 1 gramme, in suppositories, 1 dr in 7 dr of Soft Paraffin or other diluent, for an ointment

B P C employs Alcohol (45 p c) and evaporates to dryness and powders it
Official in Mex

GOSSYPIUM HAMAMELIDIS —Tincture of Hamamelis $\frac{1}{2}$ fl oz, Glycerin 10 minims, Cotton Wool, in a thin sheet, 60 grains Mix the Tincture and Glycerin, and saturate the wool evenly with the mixture Dry by exposure to the air Astringent and sedative

SUPPOSITORIUM HAMAMELIDIS —Extract of Hamamelis, $\frac{1}{2}$ grain, Oil of Theobroma, 15 grains —*Samaritan*

HAMAMELIN —A powdered extractive prepared from either the Leaves or the Bark of Hamamelis Virginiana

Dose —1 to 5 grains = 0 065 to 0 32 gramme

Two forms of Hamamelin are known in commerce, the green powder (non hygroscopic) prepared from the Leaves, and a chocolate brown hygroscopic amorphous powder prepared from the Bark

Hamamelin prepared from the Leaves with strong Alcohol was far more efficacious in suppositories than the resinoid from the Bark —*C D '98*, 1 86, *P J '01*, 11 231

HAMAMELIDIS FOLIA. **HAMAMELIS LEAVES** *B P Syn* —

WITCH HAZEL LEAVES

The Leaves, fresh and dried, of *Hamamelis Virginiana*

Foreign Pharmacopœias —Official in Austr, Belg, Fr, Jap, Mex, Norw, Span, Swed, Swiss and U S

Tests —The Leaves yield from 5 to 8 p c of ash The inclusion of an ash limit in the *B P* is stated not to be a necessity

Preparations

EXTRACTUM HAMAMELIDIS LIQUIDUM. **LIQUID EXTRACT OF HAMAMELIS**

20 of Hamamelis Leaves, percolated with Alcohol (45 p c) until exhausted, the first 17 reserved and the remainder evaporated to an Extract, which is dissolved in the first portion, and made up with Alcohol (45 p c) to 20 (1 in 1)

Dose —5 to 15 minims = 0 3 to 0 9 c c

Foreign Pharmacopœias —Official in Austr, to yield not less than 23 p c residue, Belg, to yield 23 p c residue, Fr, Jap, Norw, Span, Swed, Swiss and U S, all 1 in 1

Tests —Liquid Extract of Hamamelis has a sp gr of 1 025 to 1 050, contains about 21 p c w/v of total solids and about 32 p c w/v of Absolute Alcohol

LIQUOR HAMAMELIDIS. SOLUTION OF HAMAMELIS EXT.
HAMAMELIDIS DEST

Fresh Hamamelis Leaves, 50, Water 100, Alcohol (90 p c), 10
Macerate in a still for 24 hours, then distil one half

Oil It probably owes its virtues to the presence of a small quantity of essential

Pond's Extract and Hazeline are products distilled from Hamamelis
Official in U S

Tests.—The Liquor has a sp gr of 0.980 to 0.985, it contains
about 16 p c w/v of Absolute Alcohol

UNGUENTUM HAMAMELIDIS (HAMAMELIS OINTMENT)

Liquid Extract of Hamamelis, $\frac{1}{4}$, Hydrous Wool Fat, $2\frac{1}{4}$ (1 in 10)

Now made with Hydrous Wool Fat in place of simple Ointment

Not Official

WITCH HAZEL SNOW—Melt 2 oz of Stearic Acid and add it to a hot
solution of Glycerin 2 fl dm, Sodium Carbonate 180 grains, in Water 10 fl oz
the mixture for one hour on a water-bath, make up the volume
with Water to 10 fl oz and add Liquor Hamamelidis 10 fl oz. Transfer to a hot
mortar and agitate very thoroughly with an egg whisk. Continue agitation
till quite thick. Let stand 12 hours, stir well and bottle.—P J '06, 1 337

This has been incorporated in the B P C as follows —

Pasta Hamamelidis Syn Witch Hazel Snow or Foam—Stearic
Acid, 10, Sodium Carbonate, 1.50, Glycerin, 1.50, Solution of Hamamelis, by
weight, 50, Distilled Water, q s to produce by weight 100

Directions for preparing are the same as **Witch Hazel Snow** given above

Not Official

HELLEBORUS

CHRISTMAS ROSE

The Rhizome and Rootlets of *Helleborus Niger*, L.
It contains the glucosides Helleborein and Helleborin.—J C S Abs '98, 1 89
(It may be noted that 'White Hellebore' is Veratrum Album, and 'Green
Hellebore' is Veratrum Viride)

Medicinal Properties—A hydragogue cathartic and emmenagogue.
Poisonous in large doses, producing gastro-intestinal inflammation

Foreign Pharmacopœias—Official in Mex (Eleboro) and Port. Not in
the others

TINCTURA HELLEBORI—Hellebore Root, 1, percolated with Alcohol
(60 p c) to yield 8 (1 in 8)

Dose—20 to 60 minims = 1.2 to 3 f 3 c c in Water

This has been incorporated in the B P C

Official in Port, 1 in 5

HEMIDESMI RADIX.

The dried Root of *Hemidesmus Indicus*, R. Brown

Imported from India

It was brought to England by Dr Ashburner about the year 1830, and was

prescribed for the same purposes as Sarsaparilla, but it did not prove satisfactory, and is now used chiefly as a flavouring agent

Official Preparation — Syrupus Hemidesmi

Descriptive Notes — Hemidesmus Root occurs in pieces about 6 inches (15 cm) or more in length and $\frac{1}{8}$ to $\frac{1}{2}$ inch (3 to 12 mm) in thickness, rarely exceeding $\frac{1}{2}$ inch (6 mm) in diameter — *B P* It is cylindrical, slightly tortuous, and longitudinally furrowed, and has transverse fissures, and is of a reddish or dark brown colour, often with a violet grey hue. On one side the cork is frequently separated and raised above the cortex. The roots are furnished with a few slender rootlets, and at the upper end with slender woody stems $\frac{1}{10}$ inch (2.5 mm) or less thick, bearing opposite leaf scars. The root has a characteristic odour resembling that of Coumarin. Laticiferous vessels are found in the cortex, the wood is yellowish and porous, showing radiate medullary rays only in the smaller pieces, in the larger pieces the rays are visible only in the longitudinal or tangential section.

Tests — It yields from 3 to 4 p.c. of ash

Preparation

SYRUPUS HEMIDESMI SYRUP OF HEMIDESMUS

Infuse 4 of Hemidesmus Root in 20 of boiling Distilled Water for 4 hours, strain, and after standing, decant the clear fluid, in which dissolve 28 of Refined Sugar with a gentle heat. It should weigh 42 (about 1 in 8)

Dose — $\frac{1}{2}$ to 1 fl. dr. = 1.8 to 3.6 cc

HIRUDO.

THE LEECH

FR., SANGSUE, GER., B. LUTEGEL, ITAL., SANGUISUGA, SPAN.,
SANGUISUELA

- 1 *Sangsuga medicinalis*, the Speckled Leech, and
- 2 *Sangsuga officinalis*, the Green Leech
- 3 *Hirudo quinquestrata*, the Five Striped or Australian Leech, is official in the *Ind* and *Col A* and for the Australian Colonies

Leeches are imported chiefly from Hamburg. They are also collected in large numbers in Spain, France, Italy and Hungary.

Used for the abstraction of blood from congested parts, in pleurisy, typhlitis, pericarditis, and in cardiac distress.

When about to apply a Leech, it should be handled as little as possible, and the part of their body should be clean, and free from grease or soap, and, if a hairy part, it should be first shaved. Several suggestions have been made, in case the Leech should refuse to bite to smear the part with Milk, or Cream, or Sugar, to apply a sinapism and thoroughly clean the part afterwards, to scratch the part with a needle. When the Leech is required to bite a particular spot, it is useful to cut a small hole in blotting paper, and place it on the part

When applying a Leech to one of the orifices of the body, the Leech should be confined in a Leech glass. Should a Leech be swallowed, a strong solution of common salt (Sodium Chloride) should be drunk.

Bleeding from Leech bites is sometimes difficult to stop. The following remedies have been applied with advantage—Matico, Solution of Ferric Chloride, Silver Nitrate Point, saturated Solution of Alum, and pressure on the part.

Foreign Pharmacopœias—Official in Belg., Dan., Dutch, Fr. (Sanguis), Ger., Hung., Ital., Jap. (Hirudines), Port. (Sanguisugas), Swed. and Swiss. Not in the others.

Descriptive Notes.—There are two species met with in European commerce, viz., the Speckled or German Leech (*Sanguisuga medicinalis*, Savigny) and the Green or Hungarian Leech (*S. officinalis*, Savigny), the former having the ventral surface greenish-yellow, spotted with black, and the latter the ventral surface olive green and not spotted with black. Leeches should weigh 1 to 5 grammes only. In the Australian Colonies, the Flat-Sipped or Australian Leech, *Hirudo guttator*, Schmarda, may be substituted for the European Leeches. It has a brown dorsal surface with five longitudinal stripes, and a greenish-yellow ventral surface not spotted. Leeches should be kept in Distilled Water with a piece of charcoal in it and in the shade. After feeding, if placed in Camphor Water they will vomit the blood they have sucked, and can then be placed in clear Distilled Water, and will be ready for use again in about 10 days. The Water requires changing about once a week.

HOMATROPINÆ HYDROBROMIDUM.

HOMATROPINE HYDROBROMIDE

HYDROBROMATE OF HOMATROPINE—B.P. Add '90

$C_{16}H_{21}NO$, HBr, d_4^{20} 1.353.49.

Fr., BROMHYDRATE D'HOMATROPINE, GEL., Ital., BROMIDRATO DI HOMATROPINA.

Colourless, small, rhombic prisms, or a white crystalline, odourless powder. It is the Hydrobromide of Homatropine Mandelic Acid Ester, which is a lower homologue of Atropine.

It possesses a bitter taste.

It should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from the light.

Solubility.—1 in 6 of Water, 1 in 18 of Alcohol (90 p.c.), insoluble in Ether and in Chloroform.

Medicinal Properties.—Mydriatic. Dilates the pupil as rapidly, though not so energetically as Atropine, but its effects disappear much sooner—in about a quarter of the time. When used with Cocaine the action is quicker and more powerful.

When an oily solution is required, the alkaloid (not the salt) is dissolved in Castor Oil.

For hypodermic injection 4 grains of Homatropine Hydrobromide dissolved in 1 fl oz of sterilised Distilled Water, 6 minims = $\frac{1}{20}$ grain
 1 to 2 drops of a 1 p.c. solution in some cases of muscular asthenopia —
B M J '99, ii. 765

Dose — $\frac{1}{80}$ to $\frac{1}{10}$ grain = 0.0008 to 0.0032 gramme

Ph Ger maximum single dose, 0.001 gramme, maximum daily dose, 0.008 gramme

Official Preparation — Lamellæ Homatropinæ

Not Official — Guttes Homatropinæ, Guttes Homatropinæ cum Cocainæ, Lamellæ Homatropinæ cum Cocainæ, Homatropina, Oleum Homatropinæ cum Cocainæ

Foreign Pharmacopœias — Official in Dutch, Ger, Ital, Jap, Swed, Swiss and U S

Tests. — Homatropine Hydrobromide melts at 209° to 212° C (408.2° to 413.6° F). Neither the *BP* nor the *PG* includes a m.p., the *USP* gives 213.8° C (417° F). Its solution should be neutral in reaction towards Litmus paper. It exerts a powerful mydriatic action on the pupil of the eye. The 1 in 50 aqueous solution yields with Iodine Solution a brown precipitate, with Mercuric Chloride Test-solution a white precipitate, with Potassium Hydroxide Solution a white precipitate soluble in excess of the reagent, but no precipitate with Tannic Acid Solution or with Platonic Chloride Solution. The *PG* states that it also, after the addition of Hydrochloric Acid, yields no precipitate with Platonic Chloride Solution. The solution yields with Silver Nitrate Solution a yellowish curdy precipitate, readily soluble in Potassium Cyanide Solution, practically insoluble in Ammonia Solution, and insoluble in Nitric Acid. 1 cc of a 10 p.c. solution when cautiously mixed with Chlorine Water yields a brownish colour to Chloroform when shaken with one-fifth its volume of the latter fluid, the *USP* uses twice the volume of Chloroform. The crystalline alkaloid obtained by adding an excess of Potassium Hydroxide Solution to an aqueous solution of the salt and extracting with Ether (allowing the Ether to evaporate spontaneously), should possess a m.p. of 96° C (204.8° F). 1 centigramme of the salt mixed with a few drops of Fuming Nitric Acid, and evaporated to dryness on a water-bath, leaves a yellowish residue, which, when cool, yields on the addition of a freshly-prepared Alcoholic Potassium Hydroxide Solution a reddish-violet colour. The *USP* states that the salt yields an evanescent pink colour, changing rapidly to green when mixed with Sulphuric Acid containing a crystal of Potassium Bichromate.

The more generally occurring impurities are alkaloids other than Homatropine (Atropine, Hyoscyamine, and Hyoscine), and mineral matter. The *BP* states that a 2 p.c. aqueous solution yields no precipitate on the cautious addition of Ammonia Solution previously diluted with twice its volume of Water. A 5 p.c. solution of Atropine Sulphate with Ammonia Solution under these conditions gives a distinct turbidity, but with Hyoscyamine and Hyoscine Hydrobromides no reaction is visible. A 1 p.c. solution of Atropine Sulphate remains unchanged by the Ammonia and Mercuric Chloride Test described below. Any

salt of Atropine or Hyoscyamine under exactly similar conditions will give the same reaction but with Hyoscine no formation of Mercuric Oxide appears to take place. The most characteristic test for Homatropine is that described above, with fuming Nitric Acid and Alcoholic Potassium Hydroxide Solution. It distinguishes it from Atropine, the latter giving a deep purple coloration, as does also Hyoscyamine and Hyoscine, but in the case of the two latter, the coloration is less intense and more transient. It should leave no weighable residue when ignited with free access of air, indicating mineral impurity.

Ammonia and Mercuric Chloride—If 1 cc of a 1 in 100 aqueous solution of the salt (0.01 grammes of the salt) be made alkaline with ammonia water, shaken out with Chloroform, the chloroform evaporated to dryness, the residue should turn yellow on warming with about 1.5 cc of a solution made by dissolving 1 volume of Chlorine in 50 parts of a mixture of Alcohol, 5 parts of Atropine and Hyoscyamine, indicating the absence of most other alkaloids except Atropine and Hyoscyamine.

Platinic Chloride—Aqueous solution of the salt is not precipitated by TS of Platinic Chloride, after the addition of Hydrochloric Acid, *P. G.*

Nitric Acid and Alcoholic Potassium Hydroxide Solution—If 0.01 grammes of the salt be added to 5 drops of Nitric Acid and evaporated to dryness in a porcelain dish, the residue should not become a violet colour upon the addition of a few drops of Alcoholic TS of Potassium Hydroxide, *U. S. P.* 0.01 grammes evaporated with 5 drops of Fuming Nitric Acid in a porcelain dish on cooling and adding Alcoholic Solution of Potassium Hydroxide, assumes a transient violet colour, quickly becoming reddish-yellow.

Preparation.

LAMELLÆ HOMATROPINÆ DISCS OF HOMATROPINE

Discs of Gelatin and Glycerin, each weighing about $\frac{1}{10}$ grain = 0.0013 grammes, and containing 0.00065 grammes of Homatropine Hydrobromide.

Not Official.

GUTTÆ HOMATROPINÆ—Homatropine Hydrobromide, 4 grains, Distilled Water, 1 fl. oz.—*London Ophthalmic and Guy's*

GUTTÆ HOMATROPINÆ CUM COCAINA—Homatropine Hydrobromide, 4 grains, Cocaine Hydrochloride, 10 grains, Distilled Water, 1 fl. oz.—*London Ophthalmic and Westminster Ophthalmic*

LAMELLÆ HOMATROPINÆ CUM COCAINA—Each disc contains $\frac{1}{10}$ grain of Homatropine Hydrobromide and $\frac{1}{10}$ grain of Cocaine Hydrochloride.—*London Ophthalmic*

HOMATROPINA—Colourless crystals, not deliquescent, nearly insoluble in Water, but soluble 1 in 80 of Olive Oil, 1 in 20 of Castor Oil. They combine readily with Oleic Acid. An ointment is required.

Used in cases where an oily preparation is required.

Foreign Pharmacopœias—Official.

Homatropine Hydrochloride and **Homatropine Hydrobromide**—Both salts in white crystalline powders. Both salts are readily soluble in Water, and in Alcohol (90 per cent).

OLEUM HOMATROPINÆ CUM COCAINA—Homatropine, pure, 10 grains, Cocaine (alkaloid), 10 grains, Castor Oil, 1 fl. oz. Heat together till dissolved.—*London Ophthalmic*

salt of Atropine or Hyoscyamine under exactly similar conditions will give the same reaction, but with Hyoscyamine no formation of Mercuric Oxide appears to take place. The most characteristic test for Homatropine is that described above, with Fuming Nitric Acid and Alcoholic Potassium Hydroxide Solution. It distinguishes it from Atropine, the latter giving a deep purple coloration, as do also Hyoscyamine and Hyoscyamine, but in the case of the two latter, the coloration is less intense and more transient. It should leave no weighable residue when ignited with free access of air, any residue indicating mineral impurity.

Ammonia and Mercuric Chloride—If 1 cc of a 1 in 100 aqueous solution of the salt (0.01 gramme of the Salt, *U S P*) be made alkaline with Ammonia Water, shaken out with Chloroform, and the chloroformic solution evaporated to dryness, the residue should turn yellow, and finally brick red, when warmed with about 1.5 cc of a solution made by dissolving 1 part of Mercuric Chloride in 50 parts of a mixture of Alcohol, 5 volumes, and Water, 3 volumes; indicating the absence of most other alkaloids except Atropine and Hyoscyamine, *U S P* and *B P*.

Platinic Chloride—An aqueous solution of the salt is not precipitated by TS of Platinic Chloride, *U S P*, after the addition of Hydrochloric Acid, *P G*.

Nitric Acid and Alcoholic Potassium Hydroxide Solution—If 0.01 gramme of the salt be added to 5 drops of Nitric Acid and evaporated to dryness in a porcelain dish, the residue should not acquire a violet colour upon the addition of a few drops of Alcoholic TS of Potassium Hydroxide, *U S P*, 0.01 gramme evaporated with 5 drops of Fuming Nitric Acid in a porcelain dish on cooling and adding Alcoholic Solution of Potassium Hydroxide, assumes a transient violet colour, quickly becoming reddish yellow.

Preparation

LAMELLÆ HOMATROPINÆ Discs of HOMATROPINE
Discs of Gelatin and Glycerin, each weighing about $\frac{1}{10}$ grain =
0.0013 gramme, and containing $\frac{1}{100}$ grain = 0.00065 gramme of
Homatropine Hydrobromide

Not Official

GUTTÆ HOMATROPINÆ—Homatropine Hydrobromide, 4 grains, Distilled Water, 1 fl oz.—*London Ophthalmic and Guy's*

GUTTÆ HOMATROPINÆ CUM COCAINA—Homatropine Hydrobromide, 4 grains, Cocaine Hydrochloride, 10 grains, Distilled Water, 1 fl oz.—*London Ophthalmic*

Homatropine Hydrobromide, 7 grains, Cocaine Hydrochloride, 10 grains, Boric Acid, 5 grains, Distilled Water, 1 fl oz.—*Westminster Ophthalmic*

LAMELLÆ HOMATROPINÆ CUM COCAINA—Each disc contains $\frac{1}{10}$ grain of Homatropine Hydrobromide, and $\frac{1}{10}$ grain of Cocaine Hydrochloride.—*London Ophthalmic*

HOMATROPINA—Colourless crystals, not deliquescent, nearly insoluble in Water, but soluble 1 in 80 of Olive Oil, 1 in 20 of Castor Oil. They combine readily with Oleic Acid.

Used in cases where an oily preparation or an ointment is required.

Foreign Pharmacopœias—Official in Mex.

Homatropine Hydrochloride and Sesquicyclate form colourless crystals or white crystalline powders. Both salts are readily soluble in Water, and in Alcohol (90 p c.).

OLEUM HOMATROPINÆ CUM COCAINA—Homatropine, pure, 10 grains, Cocaine (alkaloid), 10 grains, Castor Oil, 1 fl oz. Heat together till dissolved.—*London Ophthalmic*.

salt of Atropine or Hyoscyamine under exactly similar conditions will give the same reaction, but with Hyoscyamine no formation of Mercuric Oxide appears to take place. The most characteristic test for Homatropine is that of the reaction with Fuming Nitric Acid, and from Atropine, Potassium Hydroxide Solution. It distinguishes it from Atropine, the latter giving a deep purple coloration, and also Hyoscyamine and Hyoscyne, but in the case of the two latter, the coloration is less intense and more transient. It should leave no weighable residue when ignited with free access of air, indicating mineral impurity.

Ammonia and Mercuric Chloride — A solution of the salt (0.01 gramme) in 10 c.c. of Ammonia Water, shaken out with 10 c.c. of Potassium Hydroxide Solution, and finally brick-red, warmed with about 15 c.c. of a solution made by dissolving 1 part of Mercuric Chloride in 50 parts of a mixture of Alcohol, 5 volumes, and Water, 3 volumes, indicating the presence of most other alkaloids except Atropine and Hyoscyamine.

Platinic Chloride — An aqueous solution of the salt is not precipitated by TS of Platinic Chloride, *USP*, at 60°C.

Nitric Acid and Alcoholic Potassium Hydroxide — If 0.01 gramme of the salt be added to 5 drops of Nitric Acid in a porcelain dish, the residue should not acquire a violet color upon the addition of a few drops of Alcoholic TS of Potassium Hydroxide, *USP*, 0.01 gramme evaporated with 5 drops of Fuming Nitric Acid in a porcelain dish on cooling and adding Alcoholic water-bath leaves a faint yellow residue which, on cooling and adding Alcoholic Solution of Potassium Hydroxide, assumes a transient violet colour, quickly becoming reddish-yellow.

Preparation

LAMELLÆ HOMATROPINÆ

Discs of Gelatin and Glycerin, each weighing about $\frac{1}{10}$ grain = 0.0013 gramme, and containing $\frac{1}{100}$ grain = 0.00065 gramme of Homatropine Hydrobromide.

Not Official

GUTTÆ HOMATROPINÆ — Homatropine Hydrobromide, 4 grains, Distilled Water, 1 fl oz. — *London Ophthalmic*

GUTTÆ HOMATROPINÆ CUM COCAINA — Homatropine Hydrobromide, 4 grains, Cocaine Hydrochloride, 10 grains, Distilled Water, 1 fl oz. — *London Ophthalmic*

GUTTÆ HOMATROPINÆ CUM BORIC ACID — Homatropine Hydrobromide, 7 grains, Boric Acid, 5 grains, Distilled Water, 1 fl oz. — *Manchester Ophthalmic*

LAMELLÆ HOMATROPINÆ CUM COCAINA — Each disc contains $\frac{1}{10}$ grain of Homatropine Hydrobromide, and $\frac{1}{10}$ grain of Cocaine Hydrochloride. — *London Ophthalmic*

HOMATROPINA — Colourless crystals, not deliquescent, nearly insoluble in Water, but soluble 1 in 80 of Olive Oil, 1 in 20 of Castor Oil. They combine readily with Oleic Acid.

Used in cases where an oily preparation or an ointment is required.

Foreign Pharmacopœias — Official

Homatropine Hydrochloride and Cocaine Hydrochloride — Both salts are white crystalline powders. Both salts are readily soluble in Water, and in Alcohol (90 p.c.).

OLEUM HOMATROPINÆ CUM COCAINA — Homatropine, pure, 10 grains, Cocaine (alkaloid), 10 grains, Castor Oil, 1 fl oz. Heat together till dissolved. — *London Ophthalmic*

Not Official

HORDEUM DECORTICATUM

PEARL BARLEY

The dried Seed of *Hordeum distichum*, L. divested of its early integuments, from plants cultivated in Britain

Foreign Pharmacopœias—Official in Fr (Orge Perlé), Port (Cevada Santa), Mex and Span (Cebada). Not in the others

DECOCTUM HORDEI—Pearl Barley, 1, wash the Barley with cold Water, and reject the washings, boil the washed Barley with 15 of Distilled Water for 20 minutes in a covered vessel, and strain. Product about 10 (about 1 in 10)

This has been incorporated in the *B P C*

Foreign Pharmacopœias—Official in Fr (Tisane d'Orge), 1 in 50. Not in the others. Decoctum Hordei Compositum, 1 in 50 is official in Span

Medicinal Properties—Nutritive and demulcent, used in catarrhal conditions of the respiratory and urinary systems, as a drink in febrile diseases, and to dilute cow's Milk for feeding children, thus forming a more easily digested curd

Dose—1 to 4 fl oz = 28 to 113 c c

DECOCTUM HORDEI TARTARISATUM—Acid. Potassium Tartrate, 80 grains, the Peel of $\frac{1}{2}$ Lemon, Sugar, 2½ oz, Decoction of Barley, 40 oz, boil and strain—*St Georg's*

HYDRARGYRUM.

MERCURY

Hg, eq 198 80

Fr, MERCUR PURIFIÉ, GER, QUECKSILBER, ITAL, MERCURIO, SPAN, MERCURIO

A shining, silver-white, metallic-looking fluid obtained from native Mercuric Sulphide

It should be kept in strong, well closed bottles

Solubility—Insoluble in the usual solvents, insoluble in Hydrochloric Acid, insoluble in cold Sulphuric Acid, but dissolved by hot Sulphuric Acid with evolution of Sulphur Dioxide. It dissolves readily and completely in Nitric Acid

Medicinal Properties—Mercury as a metal is seldom given alone. In a state of minute subdivision with Chalk, or in pill form, however, it has the effect of increasing the various secretions, and is itself absorbed by all the tissues of the body. It is an alterative, indirect cholagogue, purgative, diuretic, and a glandular stimulant. When given as a purgative it is usually combined with other purgatives, or followed by a purgative saline

Of great use, internally, in primary and secondary, and with Iodides in tertiary syphilis, but the doses should not be such as to cause salivation

Externally, by means of the ointment, oleate or liniment, in syphilis, in parasitic skin diseases, and as a stimulant in chronic

synovitis, peritonitis and other chronic inflammations, and glandular enlargements

See also under the various salts of Mercury

Two cases of acute intestinal obstruction successfully treated with Quick-silver — *B M J* '02, i 1023

Of the drugs frequently used in the treatment of syphilis, Blue Ointment is regarded as of most value — *L* '01, ii 1038

As an inunction ($\frac{1}{2}$ to 1 drm of the ointment well rubbed in at night before bedtime) it forms one of the most satisfactory ways of exhibiting Mercury — *B M J* '00, ii 1762

A mercurial cream prepared with a Lanolin ointment, and containing Carbolic Acid for use as an intramuscular injection in the treatment of syphilis — *B M J* '03, i 1258

Metallic Mercury still continues to be largely used in the treatment of syphilis, and preparations for use as inunctions or for intramuscular injection are in good demand. For intramuscular injection the treatment of syphilis a preparation made according to the following formula is stated (*M P* '06, i 149) to be useful — Purified Mercury, 40 grammes, sterilised Lanolin, 12 grammes, white sterilised Vaseline, 12 grammes, sterilised liquid Vaseline, 35 grammes. One c.c. contains $\frac{1}{2}$ gramme Mercury. The average dose is 7 or 8 centigrammes.

The administration of Mercury internally is stated (*B M J* '05, i 700) to be specially apt to cause symptoms of poisoning when combined with the extensive use of Tar externally.

Of the numerous salts and preparations of Mercury which have from time to time been recommended in the treatment of syphilis, attention still seems to centre round those preparations partaking of the nature of an ointment, and which can be used by inunction. In the *L* '04, i 1405, 15 grains Ung Hydrarg are recommended to be gently rubbed over the abdomen or the inside of the thigh or arm at night, and then covered with a flannel bandage until the following morning, when it is washed off, and this treatment is repeated unless the skin shows signs of irritation.

0.03 gm. of a mixture of two parts of metallic Mercury one part of Lanolin and liquid Paraffin, has been recommended (*B M J* '04, ii 1702) for subcutaneous injection. Intramuscular or intravenous injections unsuitable in infants, owing to the pain and the risk of inflammation (*L* '04, ii 1405).

A cream containing Mercury, Lanolin and Carbolic Soft Paraffin (white) is used in the Royal Navy — *B M J* '07, ii 512.

Official Preparations — Emplastrum Ammoniaci cum Hydrargyri, Emplastrum Hydrargyri, Hydrargyrum cum Creta, Liquor Hydrargyri Nitratis Acidus, Linimentum Hydrargyri, Pilula Hydrargyri, Unguentum Hydrargyri, Unguentum Hydrargyri Compositum, and Unguentum Hydrargyri Nitratis.

Not Official — Mercurial Cream (Squire), Mercury Plaster Mull, Mercury and Carbolic Plaster Mull, Oleum Cinereum Parogenum Hydrargyri, Pilula Hydrargyri Carbolicæ, Pilula Hydrargyri cum Creta, Pilula Hydrargyri cum Rheo, Suppositoria Hydrargyri, Unguentum Hydrargyri Mitius, Unguentum Cinereum, Vasolimentum Hydrargyri, Hyrgolum, Hydrargyri Benzoes, Hydrargyrum Carbolicum, Hydrargyrol, Hermophenyl Hydrargyri Cyanidum, Injectio Hydrargyri Cyanidi, Mercury Zinco-Cyanide, Unguentum Hydrargyri et Zinci Cyanidi, Hydrargyri Ethylenediamine Citras, Hydrargyri Gallicæ, Hydrargyri-Naphthol-acetas, Hydrargyri Salicylas, Hydrargyri Succimidum, Hydrargyri Sulfides, Unguentum Hydrargyri Sulphatis Flavæ, Hydrargyri Tannas, Hydrargyri Thymolacetas.

Foreign Pharmacopœias — Official in all

Tests — Mercury has a sp. gr. of 3.5. It solidifies at -39.4°C (-39°F). It boils at 360°C (680°F), and volatilises slightly even at ordinary temperatures. The fully reduced solution in Nitric Acid, freed from excess of Nitric Acid, adds with Ammonia Solution a

white precipitate, with Potassium or Sodium Hydroxide a yellow precipitate, with Potassium Iodide Solution a bright scarlet precipitate, soluble in excess of the reagent and in a considerable excess of the Mercuric salt, excess of Hydrogen Sulphide yields a black precipitate insoluble in Ammonium Hydrosulphide Solution, and in hot diluted Nitric Acid Solution. A bright piece of Copper foil immersed in the solution is coated with a grey film which, on rubbing, shows a bright silvery lustre. When the coated foil is heated in a dry clean test-tube the Mercury condenses on the sides of the tube in minute globules. The solution yields with Stannous Chloride Solution first a greyish-white precipitate of Mercurous salt and subsequently a grey precipitate of metallic Mercury. The *U S P* requires that it shall contain not less than 99.9 p.c. of metallic Mercury, but gives no method of determination. It also requires that globules of Mercury dropped upon white paper should roll about freely and leave no streaks or traces. It should present a bright surface even after agitation in contact with air.

The more generally occurring impurities are foreign metals and fixed residue. More than slight traces of foreign metals may be detected by the Sodium Thiosulphate Test described below, fixed residue by the volatilisation test.

Sodium Thiosulphate—On boiling 5 grammes of Mercury with 5 c.c. of Water and 4.5 grammes of Sodium Thiosulphate in a test tube for about one minute, the Mercury should not lose its lustre, and should not acquire more than a slightly yellowish shade, indicating the absence of more than slight traces of foreign metals, *U S P*.

Volatilisation—At a temperature of 357.25° C (675.05° F) it is completely volatilised, leaving no appreciable residue, *U S P*; it is volatilised leaving an insignificant amount of fixed residue below visible redness, *B P*.

Preparations

EMPLASTRUM HYDRARGYRI **MERCURIAL PLASTER.**

3 oz (by weight) of Mercury is rubbed with a heated mixture of 56 grains of Olive Oil, and 8 grains of Sublimed Sulphur, and finally incorporated with 6 oz of melted Lead Plaster (about 1 in 3).

Foreign Pharmacopœias—Official in Austr, Belg, Fr, Ger, Hung, Ital, Jap, Norw, Russ and Swed, 1 in 5, Dan and U S, 8 in 10, Dutch, 1 in 4, Mex, 1 in 5.57, Span, 1 in 7.5, Swed, 1 in 3. The ingredients differ considerably.

EMPLASTRUM AMMONIACI CUM HYDRARGYRO **AMMONIACUM AND MERCURY PLASTER**

3 oz (by weight) of Mercury treated as above with Olive Oil and Sulphur and mixed with 12 oz of purified Ammoniacum (nearly 1 in 5).

Applied in glandular swelling in chronic hepatic enlargement, syphilitic nodes, and in chronic synovitis.

Foreign Pharmacopœias—Official in U S, resembles Brit. Not in the others.

LINIMENTUM HYDRARGYRI **LINIMENT OF MERCURY**

Mix 1 oz of Mercury Ointment with Liniment of Camphor to

make $1\frac{1}{2}$ fl oz, and add 160 minims of strong solution of Ammonia diluted with Liniment of Camphor to $1\frac{1}{2}$ fl oz

(1 Ointment in 3, or 1 of Mercury in 6)

A stimulating Liniment, applied as an absorbent to swollen joints, or placed with Lint in the arm pits, or rubbed into the abdominal wall in tubercular peritonitis

PILULA HYDRARGYRI MERCURY PILL *B P Syn* — **BLUE PILL**

2 (by weight) of Mercury intimately mixed with 3 of Confection of Roses, and finally with 1 of powdered Liquorice Root (1 in 3)

8 commercial samples examined contained 28 to 41 p.c. of Mercury, and little or no Oxide, 5 of the 8 samples were prepared with Confection of Hips — *P J* (3) xv 230

Dose — 4 to 8 grains = 0.26 to 0.52 gramme

Foreign Pharmacopœias — Official in Fr, *Pilules Mercurielles*, *Jap* *Mix* *de* *Azules*, Port, *Pilulas Mercuriales*, Swed, *Pilulæ* *Mercuriales*, *U S*, *Mix* *de* *Hydrargyri*, all 1 in 3. Not in the others

UNGUENTUM HYDRARGYRI MERCURY OINTMENT

Mercury (by weight), 16, Lard, 16, Prepared Suet, 1

(nearly 1 in 2)

Official Preparations. — Used in the preparation of Linimentum Hydrargyri and Unguentum Hydrargyri Compositum

Foreign Pharmacopœias — Official in Austr, Belg, Dan, *Pomada Mercurialis*, Span (*Pomada Mercurialis*), 3 in 10, Fr, Ital (*Pomata Mercurialis*), Mex (*Unguento de Mercurio Doble*), Port and US, 1 in 2, Fr has also *Pomade Mercurielle Faible*, 1 in 8, Ger, Hung, Jap, Russ, Swed and Swiss (*Ung Hydr Ciner*), 1 in 3. Span has also *Pomada Mercurialis Simple*, 3 in 20

The Brussels Conference adopted a strength of 30 p.c. for Unguentum Hydrargyri

Mercury, 50, Oleate of Mercury, 2, Suet, 3, Benzoinated Lard, 25 — *US*

UNGUENTUM HYDRARGYRI COMPOSITUM COMPOUND MERCURY OINTMENT

Mercury Ointment, 10, Yellow Beeswax, 6, Olive Oil (by weight), 6, Camphor, in flowers, 3. Mix the Beeswax, Olive Oil, and Mercury Ointment with the aid of heat, add the Camphor, triturate until cold (1 Mercury in 5)

Contains rather less Mercury Ointment than B P '65, and therefore its action is modified, as previously suggested in the *Companion*.

This is Scott's celebrated absorbent Ointment (Scott's dressing), the Soap Cerate being replaced by the Oil and Beeswax

It is an admirable Ointment to apply to chronic joint enlargement.

Not Official

MERCURY PLASTER MULL (*Unna*) — Containing 1 grain = 0.06 grammes of Mercury to the square inch

MERCURY AND CARBOLIC PLASTER MULL (*Unna*) — Containing 1 grain = 0.06 gramme of Mercury and $\frac{1}{2}$ grain = 0.02 gramme of Carbolic Acid to the square inch

MERCURIAL CREAM (*Scott*) — Boiled re-distilled Mercury, by weight, 48 grains, sterilised anhydrous Lanthanum, by weight, 240 grains, pure sterilised Olive Oil, 7 s to produce 1 fl oz.

10 minims = 1 grain of pure mercury

Dose.—10 minims = 0.6 c.c. by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 5 parts of carbolic oil of 2% strength, 10 minims of the resultant grey cream contained 1 grain of metallic Mercury. Lang's formula, published in 1888, suggested the original principle of the process, namely the minute subdivision of the metallic Mercury by means of Lanolin, and the thinning of the emulsion with Olive Oil.

The following formula is given *B M J* '03, 1258 by Colonel F. J. Lambkin, R. A. M. C.—Mercury, 2 drms., anhydrous Lanolin, 2 drms. by weight, Carbolic Acid, 4 drms., Carbolic Acid, 2 p.c., by measure.

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under-mentioned formulas are given by Colonel F. J. Lambkin, R. A. M. C., in *L* '07, 114.

Mercury, 10 grammes, absolute Rosin and Camphoric Acid (Creo Camph) of each equal parts, 20 c.c., Palmitin Basis to 100 c.c.

Calomel, 5 grammes, absolute Rosin and Camphoric Acid (Creo Camph) of each equal parts, 20 c.c., Palmitin Basis to 100 c.c.

As a basis for Calomel Injection used in Syphilis, Dr Allaire ('Nouveaux Remèdes') recommends Palmitin prepared from palm oil. It does not become rancid, easily saponifies in the body, and is readily absorbed. A little Guaiacoloid (a combination in molecular proportions of Guaiacol and Camphor) is added to the injection.—*C D* '07, 1141.

OLEUM CINEREUM (Grey Oil).—White Vaseline, 25, Mercury Ointment, 1, Mercury, 19.5, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. Mercury.—*P J* (3) xix 704.

For hypodermic injection in syphilis. **Dose**—1 to 2 minims.—*B M J* '88, 1296, *T G* '94, 819.

A modification of 'Grey Oil,' is Mercury, 1, Lanolin anhydrous, 2, Carbolic Oil (2 p.c.), 1, all by weight. 10 mms. used for each injection.—*B M J* '98, 1485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, *q s* to produce (by weight) 100. **Dose**—1 to 2 minims.—*B P C*.

The following formula appears in the *Fr. Codex* (1908) under the title of

Huile Grise.—Purified Mercury, 1, anhydrous Lanolin, 26, Vaseline Oil (Huile de Vaseline medicinale), 60. The Lanolin and the Vaseline Oil are sterilised separately in glass flasks in an autoclave at 120° C (248° F) for 20 minutes. A pestle and mortar are heated by means of burning Alcohol, and placed therein are the Mercury, and the Wool Fat. The metallic particles are triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should weigh 126 grammes, and measure 100 c.c., and therefore contains almost exactly centigrammes of Mercury per c.c., or 40 p.c. w/v, and should be transferred immediately to phials of two, five, and ten c.c. capacity previously sterilised at 120° C (356° F).

PILULA HYDRARGYRI CUM OPIO.—Mercury Pill Mass, 5 grains, Opium, in powder, $\frac{1}{2}$ grain.—*St. Thomas*.

This has been incorporated in the *B*.

Mercurial Pill to 32 grains, Opium, in powder, $\frac{1}{2}$ grain.—*Guy's*.

Mercury Pill, 5 grains, Opium, in powder, $\frac{1}{2}$ grain.—*University* (No. 3) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM EO.—Mercury Pill Mass, 2½ grains, Compound Rhubarb Pill Mass, 2½ grains.—*Thomas's* and *London Ophthalmic and King's*.

This has been incorporated in the *B*.

SUPPOSITORIA HYDRARGYRI.—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

Dose—10 minims = 0.6 c.c. by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 5 parts of carbolic oil of 2 p.c. strength, 10 minims of the resultant grey cream contained 1 grain of metallic Mercury. Lang's formula, published in 1888, suggested the original principle of the process, namely the minute subdivision of the metallic Mercury by means of Lanolin, and the thinning of the emulsion with Olive Oil.

The following formula is given (*B M J* '03, 1258) by Colonel F. J. Lambkin R. A. M. C.—Mercury, 2 drms., anhydrous Lanolin, 2 drms. by weight, Paroleine 4 drms., Carbolic Acid, 2 p.c., by measure.

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under-mentioned formulas are given by Colonel F. J. Lambkin R. A. M. C., in *L* '07, 114.

Mercury, 10 grammes, and Camphoric Acid (Creo-Camph) of each equal parts, 20 c.c., 100 c.c.

Calomel, 5 grammes, absolute Creosote and Camphoric Acid (Creo Camph) of each equal parts, 20 c.c., Palmitic Basis to 100 c.c.

As a basis for Calomel in Syphilis, Dr. Allaire ('Nouveaux Remèdes') recommends Palm Oil. It does not become rancid, easily saponifies in the body, and is readily absorbed. A little Guaiacaloid (a combination in 100 c.c. of Guaiacal and Camphor) is added to the injection—*C D* '07, 114.

OLEUM CINEREUM (Grey Oil)—White Vaseline, 2.5, Mercury Ointment, 1, Mercury, 19.5, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. of Mercury—*P J* (3) xix 704.

For hypodermic injection in syphilis. **Dose**—1 to 2 minims—*B M J* '88, 1250; *J G* 95.

A modification of this Oil is Mercury, 1, Tincture of Iodine, 2, Carbolic Oil (2 p.c.), and Olive Oil 10 minims used—*B M J* '98, 1485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, *q.s.* to produce (by weight) 100. **Dose**—1 to 2 minims—*B P C*.

The following is in the *Fr. Codex* (1908) under the title of

Huile Gise. Purified Mercury, 40, anhydrous Lanolin, 26 Vaseline Oil (Huile de Vaseline médicale), 60. The Lanolin and the Vaseline Oil are sterilised separately in glass flasks in an autoclave at 120° C (248° F) for 20 minutes. A pestle and mortar are sterilised by means of burning Alcohol, and placed therein are the Mercury, and then the Wool Fat. The metallic particles are triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should measure 100 c.c., and therefore contains almost exactly 40 p.c. of Mercury per c.c., or 40 p.c. w/v, and should be transferred to phials of two, five, and ten c.c. capacity previously sterilised at 120° C (248° F).

PILULA HYDRARGYRI CUM OPIO—Mercury Pill Mass, 5 grains, Opium, in powder $\frac{1}{2}$ grain—*St. Thomass*.

This has been incorporated in the *B P C*.

Mercurial Pill to $3\frac{1}{2}$ grains, Opium, in powder, $\frac{1}{2}$ grain—*Guy's*.

Mercury Pill, 5 grains, Opium, in powder, $\frac{1}{2}$ grain—*University* (No 3) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM RHEO.—Mercury Pill Mass, $2\frac{1}{2}$ grains, Compound Rhubarb Pill Mass, $2\frac{1}{2}$ grains—*St. Thomas's* and *London Ophthalmic* and *King's*.

This has been incorporated in the *B P C*.

SUPPOSITORIA HYDRARGYRI—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

make $1\frac{1}{2}$ fl oz, and add 160 minims of strong solution of Ammonia diluted with Liniment of Camphor to 1 fl oz
(1 Ointment in 3, or 1 of Mercury in 6)

A stimulating Liniment, applied is in absorbent to swollen joints, or placed with Lint in the sinu pits, or rubbed into the abdominal wall in tubercular peritonitis

PILULA HYDRARGYRI **MERCURY PILLS** *B P Sgr* — **BLU PILL**

2 (by weight) of Mercury intimately mixed with 3 of Confection of Roses, and finally with 1 of powdered Liquorice Root (1 in 3)

8 commercial samples examined contained 28 to 41 p.c. of Mercury, and little or no Oxide, 5 of the 8 samples were impure with Confection of Hips — *P J* (3) vi 230

Dose — 4 to 8 grains = 0.26 to 0.53 gramme

Foreign Pharmacopœias — Official in *B P*, *Pilulas Mercuriellæ* Simplex, *Jap*, *Mex*, *Pilulas Azules*, *Port*, *Pilulas Mercuriales*, *Swed*, *Pilula Hydrargyri*, *US*, *Massæ Hydrargyri*, all 1 in 3. Not in the others

UNGUENTUM HYDRARGYRI **MERCURY OINTMENT**

Mercury (by weight), 16, Lard, 16, Prepared Suet, 1

(nearly 1 in 2)

Official Preparations — Used in the preparation of Linimentum Hydrargyri and Unguentum Hydrargyri Compositum

Foreign Pharmacopœias — Official in *Austr*, *Belg*, *Dan*, *Den*, *Span* (*Pomada Mercurialis*), 9 in 10, *Fr*, *Ital* (*Pomata Mercurialis*), *Mex* (*Unguento de Mercurio Doble*), *Port* and *US*, 1 in 2, *Fr* also *Pomade Mercurielle Faible*, 1 in 8, *Ger*, *Hung*, *Jap*, *Russ*, *Swed* and *Swiss* (*Ung Hydr Cinei*), 1 in 3. *Span* has also *Pomada Mercurialis Simple*, 8 in 20

The Brussels Conference adopted a strength of 80 p.c. for Unguentum Hydrargyri

Mercury, 50, Oleate of Mercury, 2, Suet, 3, Benzomated Lard, 25 — *US*

UNGUENTUM HYDRARGYRI COMPOSITUM **COMPOUND MERCURY OINTMENT**

Mercury Ointment, 10, Yellow Beeswax, 6, Olive Oil (by weight), 6, Camphor, in flowers, 3. Mix the Beeswax, Olive Oil, and Mercury Ointment with the aid of heat, add the Camphor, triturate until cold
(1 Mercury in 5)

Contains rather less Mercury Ointment than *B P* '85, and the manipulation is modified, as previously suggested in the *Companion*

This is Scott's celebrated absorbent Ointment (*Scott's dressing*), the Soap Cerate being replaced by the Oil and Beeswax

It is an admirable Ointment to apply to chronic joint enlargements

Not Official

MERCURY PLASTER MULL (*Unna*) — Containing 1 grain = 0.06 gramme of Mercury to the square inch

MERCURY AND CARBOLIC PLASTER MULL (*Unna*) — Containing 1 grain = 0.06 gramme of Mercury and $\frac{3}{4}$ gr in = 0.02 gramme of Carbolic Acid to the square inch

MERCURIAL CREAM (*Squire*) — Pure re-distilled Mercury, by weight, 48 grains, sterilised anhydrous Lanolin, by weight, 240 grains, pure sterilised Olive Oil, $\frac{1}{2}$ ss to produce 1 fl oz

10 minims = 1 grain of pure metallic Mercury

Dose.—10 minims = 0.6 c.c. by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, is of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of anhydrous Lanolin, and then mixed with 5 parts of emulsified oil of 2 per cent strength, 10 minims of the resultant grey cream contained 1 grain of metallic Mercury. Lang's formula, published in 1888, is the original principle of the process, namely the minute sub-division of metallic Mercury by means of Lanolin, and the thinning of the mixture with Olive Oil.

The following formula is given (*B M J* '03, 1258) by Colonel F. J. Lambkin, R.A.M.C.—Mercury, 2 drms, anhydrous Lanolin, 2 drms by weight, Parement, 4 drms, Carbolic Acid, 2 p.c., by measure.

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under-mentioned formulas are given by Colonel F. J. Lambkin, R.A.M.C., in 1907, 1144.

Mercury, 10 grammes, absolute Croosote and Camphoric Acid (Croo-Camph.) of each equal parts, 20 c.c., Palmistia Basis to 100 c.c.

Calomel, 5 grammes, absolute Croosote and Camphoric Acid (Croo-Camph.) of each equal parts, 20 c.c., Palmistia Basis to 100 c.c.

As a basis for the preparation used in Syphilis, Dr Allaire ('Nouveaux Remèdes') recommends a mixture prepared from palm oil. It does not become rancid, easily separates from the body, and is readily absorbed. A little Guaiacoloid (a combination in molecular proportion of Guaiacol and Camphor) is added to the injection.—*C D*, '07, 1141.

OLEUM CINEREUM (Grey Oil).—White Vaseline, 2.5, Mercury Ointment, 1, Mercury, 19.5, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. of Mercury.—*P J* (3) 701.

For hypodermic injection in Syphilis. **Dose**—1 to 2 minims.—*B M J*, '88, 1296, 774.

A modification of the above Oil, is Mercury, 1, Lanolin anhydrous, 2, Carbolic Oil (2 p.c.), 1, all by weight. 10 minims used for each injection.—*B M J* '98, 1485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, 95 to produce (by weight) 100.

Dose—1 to 2 minims.—*B P C*.

The following formula appears in the *Ph Codex* (1908) under the title of

Huile Grise—Purified Mercury, 40, anhydrous Lanolin, 26, Vaseline Oil (Huile de Vaseline medicinale), 60. The Lanolin and the Vaseline Oil are sterilised separately in glass flasks in an autoclave at 120° C. (248° F.) for 20 minutes. A pestle and mortar are sterilised by means of boiling alcohol, and placed therein are the Mercury, and then the Wool Fat. The anhydrous Lanolin is triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should be transferred to phials and measure 100 c.c., and therefore contains almost exactly 40 centigrammes of Mercury per c.c., or 40 p.c. w/v, and should be transferred immediately to phials of two, five, and ten c.c. capacity previously sterilised at 180° C. (356° F.).

PILULA HYDRARGYRI CUM OPIO—Mercury Pill Mass, 5 grains, Opium, in powder, $\frac{1}{2}$ grain.—*St. Thomas's*.

This has been incorporated in the *B P C*.

Mercurial Pill to 3½ grains, Opium, in powder, $\frac{1}{2}$ grain.—*London Ophthalmic*.

Mercury Pill, 5 grains; Opium, in powder, $\frac{1}{2}$ grain.—*London Ophthalmic* (No 8) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM RHEO.—Mercury Pill Mass, 2½ grains, Compound Rhubarb Pill Mass, 2½ grains.—*St. Thomas's* and *London Ophthalmic* and *King's*.

This has been incorporated in the *B P C*.

SUPPOSITORIA HYDRARGYRI—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

Dose—10 minims = 0.6 c.c. by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 5 parts of carbolic oil of 2 p.c. strength, 10 minims of the resultant grey cream contained 1 grain of metallic Mercury. Lang's formula, published in 1888, suggested the original principle of the process, namely the minute subdivision of the metallic Mercury by means of Lanolin, and the thinning of the emulsion with Olive Oil.

The following formula is given (*B. M. J.* '03, i 125) by Colonel F. J. Lambkin, R. A. M. C.—Mercury, 2 drim, anhydrous Lanolin, 2 drim by weight, Paroleine, 4 drim, Carbolic Acid, 2 p.c., by measure.

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under-mentioned formulas are given by Colonel F. J. Lambkin, R. A. M. C., in *L.* '07, ii 11.

Mercury, 10 grammes, absolute Creosote and Camphoric Acid (Creo-Camph.) of each equal parts, 20 c.c., Palmistil Basis to 100 c.c.

Calomel, 5 grammes, absolute Creosote and Camphoric Acid (Creo-Camph.) of each equal parts, 20 c.c., Palmistil Basis to 100 c.c.

As a basis for Calomel Injection used in Syphilis, Dr Allaire ('Nouveaux Remèdes') recommends a mixture prepared from palm oil. It does not become rancid, easily saponifies in the body, and is readily absorbed. A little Guanacolid (a combination in molecular proportions of Guanacol and Camphor) is added to the injection.—*C. D.* '07, ii 111.

OLEUM CINEREUM (Grey Oil)—White Vaseline, 25, Mercury Ointment, 1, Mercury, 195, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. of Mercury.—*P. J.* (3) ix 704.

For hypodermic injection in syphilis. **Dose**—1 to 2 minims.—*B. M. J.* '88, i 1296, *T. G.* '94, 414.

A modification of Grey Oil, is Mercury, 1, Lanolin anhydrous, 2, Carbolic Oil (2 p.c.), 1, all by weight. 10 minims used for each injection.—*B. M. J.* '98, i 485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, *q. s.* to produce (by weight) 100. **Dose**—1 to 2 minims.—*B. P. C.*

The following formula appears in the *Fr. Codex* (1908) under the title of

Huile Grise—Purified Mercury, 40, anhydrous Lanolin, 26, Vaseline Oil (Huile de Vaseline medicinale), 60. The Lanolin and the Vaseline Oil are sterilised separately in glass flasks in an autoclave at 120° C (248° F) for 20 minutes. A pestle and mortar are sterilised by means of burning Alcohol, and placed therein are the Mercury, and then the Wool Fat. The metallic particles are triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should weigh 126 grammes, and measure 100 c.c., and therefore contains almost exactly 40 centigrammes of Mercury per c.c., or 40 p.c. w/v, and should be transferred immediately to phials of two, five, and ten c.c. capacity previously sterilised at 180° C (356° F).

PILULA HYDRARGYRI CUM OPIO—Mercury Pill Mass, 5 grains, Opium, in powder, $\frac{1}{2}$ grain.—*St. Thomas's*.

This has been incorporated in the *B. P. C.*

Mercurial Pill to $3\frac{1}{2}$ grains, Opium, in powder, $\frac{1}{2}$ grain.—*Guy's*.

Mercury Pill, 5 grains, Opium, in powder, $\frac{1}{2}$ grain.—*University* (No. 8) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM RHEO—Mercury Pill Mass, $2\frac{1}{2}$ grains, Compound Rhubarb Pill Mass, $2\frac{1}{2}$ grains.—*St. Thomas's* and *London Ophthalmic* and *King's*.

This has been incorporated in the *B. P. C.*

SUPPOSITORIA HYDRARGYRI.—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

Dose.—10 minims = 0.6 cc by intramuscular injection. The preparation recommended by Dr. Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 5 parts of carbolised oil of 2 per cent strength, 10 minims of the resultant given cream contained 1 grain of metallic Mercury. Lang's formula, published in 1892, the original principle of the process, namely the minute subcutaneous injection of metallic Mercury by means of Lanolin, and the thinning of the

The following formula is given by R A M C—Mercury, 2 dm, 1.1, R 71 '03, 1258) by Colonel F J Lambkin, 4 dm; Carbolic Acid, 2 p c, by 1.1, Lanolin, 2 dm by weight, Paroleine,

Dose—5 to 10 minims once a

The two under-mentioned formulas are given by Colonel F J Lambkin,
R A M C, in L '07, n 14

Mercuric Iodide 100 cc
of each equal parts 20 cc
Cresol and Camphoric Acid (Creso-Camph)
Basis to 100 cc

Calome of each equal parts 20 g. and Camphoric Acid (Creo Camph) 100 cc

As a base, for example, in the preparation of the "Remèdes" recommended by the Société de Pharmacie, Dr. Allane ("Nouveaux Remèdes") recommends the use of a mixture of 1 part of Guaiacum oil and 1 part of Camphor. It does not become rancid, easily evaporates, and is readily absorbed. A little Guaiacoloid (a combination in molecular proportions of Guaiacum and Camphor) is added to the injection — C D '07, 11 411.

OLEUM CINEREUM (Grey^{hl})—White Vaseline, 2 5, Mercury Ointment, 1, Mercury, 19 5, tincture in a warm mortar until the Mercury is incorporated, then add White Vaseline, 2 7, Liquid Vaseline, 20 All by weight

This preparation contains 40 p.c. of Mercury — *P J* (3) xix 704
For hypodermic injection in syphilis Dose — 1 to 2 minims — *B M J*

A rod injection of 1 cc of Oil, is 1, Mercury, 1, Lanolin anhydrous, 2, Carbolic Oil (2 p c), 1, all by weight 10 mgms used for each injection — *B M J* 98, 1 485

Dose — 1 to 2 minims — *B P C*

The following formula appears in the *Fr Codex* (1908) under the title of
Huile Grise—Purified Mercury:

Huile Grise - Purified Mercury,
(Huile de Vaseline mcd. 120) 60
sterilised separately in glass bottles in
minutes. A pestle and mortar are then
placed therein and the Mercury, and the
are triturated until they are extinguished
in small portions. The product should
and therefore contain almost exactly
40 per cent, and should be transferred
ten cc capacity previously sterilised at
anhydrous Jarol 26 Vaseline Oil
the Lard and the Vaseline Oil are
also Jarol 120 (240-1) for 20
used by means of burning Alcohol and
the Wool Fat. The metallic particles
and then the Liquid Paraffin is added
126 grammes, and measure 100 cc,
centigrammes of Mercury per cc, or
immediately to phials of two, five, and
10 cc (356-1)

PILULA HYDRARGYRI CUM OPIO —Mercury Pill Mass, 5 grains,
Opium, in powder, $\frac{1}{2}$ grain —*St Thomas*
This has been incorporated in the B C
Mercurial Pill to $3\frac{1}{2}$ grains — Opium in powder, $\frac{1}{2}$ grain —*Guy's*
Mercury Pill, 5 grains — Opium, in powder, $\frac{1}{2}$ grain —*University* (No 3) and
London Ophthalmic

PILULA HYDRARGYRI CUM —Mercury Pill Mass, 2½ grains,
Compound Rhubarb Pill Mass, 2½ grains,
and *King's* *St Thomas's and London Ophthalmic*

This has been incorporated in the B_C

SUPPOSITORIA HYDRARGYRI Mercury Ointment, 5 grains, Oil of
Theobroma, 10 grains, in each suppository

Dose—10 minims = 0.6 c.c. by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury, thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 5 parts of carbolic oil of 2% strength, 10 minims of the resultant grey cream contained 1 grain of metallic Mercury. Lang's formula, published in 1888, suggested the original principle of the process, namely the minute sub-division of the metallic Mercury by means of Olive Oil, and the thinning of the emulsion with Olive Oil.

The following formula is given by *B M J* '03, 1 1258 by Colonel F J Lambkin, R A M C—Mercury, 2 drms, anhydrous Lanolin, 2 drms by weight, Paroleine, 4 drms, Carbolic Acid, 2 p.c., by measure.

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under mentioned formulas are given by Colonel F J Lambkin, R A M C, in *L* '07, 11 14.

Mercury, 10 grammes absolute, Creosote and Camphoric Acid (Creo Camph) of each equal parts, 20 c.c., Palmolive Basis to 100 c.c.

Calomel, 5 grammes, absolute, Creosote and Camphoric Acid (Creo Camph) of each equal parts, 20 c.c., Palmolive Basis to 100 c.c.

As a basis for Carbolic Injection, used in Syphilis, Dr Allaire ('Nouveaux Remèdes') recommends minute quantities of palm oil. It does not become rancid, easily saponifies in the body, and is readily absorbed. A little Guaiacolid (a combination in molecular proportions of Guaiacol and Camphor) is added to the injection—*C D* '07, 11 411.

OLEUM CINEREUM (Grey Oil)—White Vaseline, 2.5, Mercury Ointment, 1, Mercury, 19.5, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 2.5, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. of Mercury—*P J* (3) xix 704. For hypodermic injection in syphilis. **Dose**—1 to 2 minims—*B M J* '88, 1 1296, *T G* '94, 319.

A modification of 'Grey Oil' is 1, Mercury, 1, Lanolin anhydrous, 2, Carbolic Oil (2 p.c.), 1, all by weight. 10 minims used for each injection—*B M J* '98, 1 485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, *q s* to produce (by weight) 100. **Dose**—1 to 2 minims—*B P C*.

The following formula appears in the *Fr Codex* (1908) under the title of

Huile Grise—Purified Mercury, 50, anhydrous Lanolin, 26, Vaseline Oil (Huile de Vaseline medicinale), 60. The Lanolin and the Vaseline Oil are sterilised separately in glass flasks in autoclave at 120° C (248° F) for 20 minutes. A pestle and mortar are sterilised by means of burning Alcohol, and then placed therein are the Mercury, and the Wool Fat. The metallic particles are triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should weigh 126 grammes, and measure 100 c.c., and therefore contains almost exactly 1 centigrammes of Mercury per c.c., or 40 p.c. w/v, and should be transferred immediately to phials of two, five, and ten c.c. capacity previously sterilised at 100° C (356° F).

PILULA HYDRARGYRI CUM OPIO—Mercury Pill Mass, 5 grains, Opium, in powder, $\frac{1}{2}$ grain—*St Thomas*. This has been incorporated in the *B C*. Mercurial Pill to 32 grains, Opium, in powder, $\frac{1}{2}$ grain—*Guy's*. Mercury Pill, 5 grains, Opium, in powder, $\frac{1}{2}$ grain—*University* (No 3) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM RHEO—Mercury Pill Mass, 2½ grains, Compound Rhubarb Pill Mass, 2½ grains—*St Thomas's* and *London Ophthalmic* and *King's*.

This has been incorporated in the *B C*.

SUPPOSITORIA HYDRARGYRI—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

Dose—10 minims = 0.6 cc by intramuscular injection. The preparation recommended by Dr Julius Althaus, in his paper before the International Medical Congress at its Berlin meeting in 1890, consisted of 1 part of metallic Mercury thoroughly rubbed up with 4 parts of purest Lanolin, and then mixed with 10 parts of carbolic oil of 2 p.c. strength, 10 minims of the resultant grey cream contained 1 gram of metallic Mercury. Lang's formula, introduced in 1888, suggested the original principle of the process, namely the absorption of the metallic Mercury by means of Lanolin, and the thinning of the emulsion with Olive Oil.

The - - - - - '03, 1258) by Colonel F. J. Lambkin
R. A. M. C., in L '07, 11 14
2 dim., Carbolic Acid 2 p.c., Lanolin, 2 dim. by weight, Paroelene,

Dose—5 to 10 minims once a week as an intramuscular injection.

The two under-mentioned formulas are given by Colonel F. J. Lambkin, R. A. M. C., in L '07, 11 14.

Mercury, 10 grammes, Carbolic Acid 2 p.c., and Camphoric Acid (Creo-Camph) of each equal parts, 20 cc., to be made up to 100 cc.

Calomel, 5 grammes, absolute Creosote and Camphoric Acid (Creo-Camph) of each equal parts, 20 cc., Palmatine Basis to 100 cc.

As a basis for Calomel Injection used in Syphilis, Dr Allaine (Remedes) recommends Palmatine prepared from palm oil. It does not saponify, easily saponifies in the body, and is readily absorbed. A little (a combination in Guaiacol and Camphor) is added to the injection—C. D. '07, 11 411.

OLEUM CINEREUM (Grey Oil)—White Vaseline, 25, Mercury Ointment, 1, Mercury, 195, triturate in a warm mortar until the Mercury is incorporated, then add White Vaseline, 7, Liquid Vaseline, 20. All by weight.

This preparation contains 40 p.c. of Mercury—P. J. (3) xix 704.

For hypodermic injection in syphilis. **Dose**—1 to 2 minims—B. M. J. '88, 1296, T. G. '94 319.

A more recent form of Grey Oil is Mercury, 1, Lanolin anhydrous 2, Carbolic Oil (2 p.c.) 10 minims used for each injection—B. M. J. 98, 1485.

Mercury, 40, Wool Fat, 10, Liquid Paraffin, 95 to produce (by weight) 100.

Dose—1 to 2 minims—B. P. C.

The following formula appears in the *Fr. Codex* (1908) under the title of

Huile Grise—Purified Mercury, 10, anhydrous Lanolin, 26, Vaseline Oil (Huile de Vaseline medicinale), 60. The Lanolin and the Vaseline Oil are sterilised separately, in glass flasks in an autoclave at 120° C (243° F) for 20 minutes. A pestle and mortar are sterilised by means of burning Alcohol and placed therein are the Mercury and the Wool Fat. The metallic particles are triturated until they are extinguished, and then the Liquid Paraffin is added in small portions. The product should weigh 126 grammes, and measure 100 cc., and therefore contains almost exactly 10 centigrammes of Mercury per cc., or 40 p.c. w/v, and should be tried in a capacity previously sterilised at 120° C (243° F).

PILULA HYDRARGYRI CUM OPIO—Mercury Pill Mass, 5 grains.

Opium in powder, $\frac{1}{2}$ grain—*St. Thomas*.

This has been incorporated in the B. C.

Mercurial Pill to 3 grains Opium 1 grain—*Guy's*.

Mercury Pill, 2 grains Opium, in 10 grains—*University* (No 3) and *London Ophthalmic*.

PILULA HYDRARGYRI CUM HEO.—Mercury Pill Mass, 2½ grains, Compound Rhubarb Pill Mass, 2½ grains—*St. Thomas* and *London Ophthalmic* and *King's*.

This has been incorporated in the B. C.

SUPPOSITORIA HYDRARGYRI—Mercury Ointment, 5 grains, Oil of Theobroma, 10 grains, in each suppository.

UNGUENTUM HYDRARGYRI MITIUS—Mercurial Ointment, 1, Lard, 2 —*P L '86*

This has been incorporated in the *B P C* under the title *Unguentum Hydrargyri Dilutum*

Mercurial Ointment (*U S P*), 67, Petrolatum, 33 —*U S P*

UNGUENTUM CINEREUM—Mercury and Lanolin, of each 1 oz., best Olive Oil, $\frac{1}{2}$ fl oz —*Look*

VASOLIMENTUM HYDRARGYRI—Mercury, 40, Wool Fat, 20, Thick Vasoliment, 60 —*Hager*

Parogenum Hydrargyri Syn Mercury Vasoliment—Mercury, 30, Wool Fat, 15, Thick Parogen, 55 —*B P C*

HYRGOLUM (Colloid Mercury)—Heavy black grains exhibiting a metallic lustre, containing 73 to 80 p c of Mercury, soluble in Water. On account of its freedom from causticity and from irritating properties, it has been suggested as an anti-syphilitic remedy in the form of a 10 p c ointment, or internally in $\frac{1}{2}$ grain dose in pill form —*L '00*, 1450, *B M J '01*, 1551

HYDRARGYRI BENZOAS $\text{Hg}(\text{C}_6\text{H}_5\text{O})_2$, eq 439.06—A white crystalline salt, practically insoluble in Water and in Alcohol (90 p c), but soluble in solutions of the Benzoates of the alkali metals. Has been used in the treatment of syphilis. The hæmostatic effects of intramuscular injections in cases of uterine hæmorrhage are stated to far surpass Ergot (*B M J '04*, 11085)

Mercury Benzoate, Bimodide and Lactate are employed (*M P '05*, 11622) in the treatment of syphilis in daily doses of $\frac{1}{2}$ grain and should be sufficiently diluted (2 c c of Water). Mercury Salicylarsenate and Hermophenyl are generally but little painful in injections in syphilis, and are given in larger doses, *e g*, $\frac{1}{2}$ to 1 grain. Hermophenyl may be employed in larger doses ($1\frac{1}{2}$ grains), but only as a weekly injection.

One centigramme ($\frac{1}{5}$ grain) Benzoate is a small daily dose in the treatment of syphilis, and 2 centigrammes daily for three weeks may be safely given (*M P '06*, 1148). A good formula is Mercury Benzoate, 1 gramme, Sodium Chloride pure, $\frac{1}{2}$ gramme, Distilled Water, 100 grammes.

Six cases of general paralysis and tabes treated by hypodermic injection of 3 centigrammes Mercury Benzoate daily for 15 days alternated by a 15 days' interval —*B M J E '02*, 1187

A suitable solution (*Desessquale and B. etonneau*) for hypodermic injection in syphilis, Mercuric Benzoate, 0.3 gramme, Ammonium Benzoate, 1.5 grammes, Sterilised Distilled Water, to 30 c c —*P J '02*, 1173

Foreign Pharmacopœias—Official in Fr

Tests—Mercuric Benzoate Solution yields the tests distinctive of Mercury given under that substance. With Ferric Chloride T.S. it yields a buff coloured precipitate. When shaken with Water and filtered, the filtrate, when acidified with Nitric Acid, yields no precipitate or turbidity with Silver Nitrate Solution. Another portion of the filtrate, when mixed with an equal volume of Sulphuric Acid, keeping the mixture cool, should yield no brown ring at the junction of the two fluids on the careful addition of Ferrous Sulphate Solution. The Benzoic Acid obtained from the salt should possess the m.p., answer the tests distinctive of Benzoic Acid, and be free from the impurities mentioned under Acidum Benzoicum. 0.5 gramme ignited with free access of air should leave no weighable residue. It contains theoretically 45.3 p c of metallic Mercury.

HYDRARGYRUM CARBOLICUM (Mercury Carbolate, Mercury Phenate) (*Schadek*)—Colourless crystals, or a white powder. Obtained by precipitating an alcoholic Solution of Mercuric Chloride with an alcoholic Solution of Phenol and Potassium Hydroxide, and evaporating nearly to dryness, with subsequent washings.

Nearly insoluble in Water, and soluble with difficulty in cold Alcohol.

Medicinal Properties—Recommended in secondary syphilis —*L '87*, 1943, *L '87*, 11277, *P J (3)* xviii 695.

Dose— $\frac{1}{2}$ to $\frac{1}{4}$ grain = 0.02 to 0.05 gr. 3 or 4 times a day in pill, also hypodermically suspended in M.

Pilula Hydrargyri Carbolicæ—Mercury Carbolate, $\frac{1}{2}$ grain, Extract of Liquorice, 1 grain, Powdered Liquorice, 1 grain, in each pill.

Dose—Two to four pills daily

Hydrargyrol (Mercury Phenol-para-sulphonate)—Brownish-red crystalline scales or crusts. Decomposed by Water with the formation of basic salts. Insoluble in Alcohol (90 p c). Introduced as an antiseptic.

A combination of the above salt with Ammonium Tartrate is known under the name of 'Asterol,' a white or reddish-white micro-crystalline powder, soluble in Water. Introduced as an antiseptic, used in the form of 2 to 5 p c solution.—*B.M.J.E.* '01, ii 64, *P.J.* '99, i 538, '99, ii 216, *C.D.* '01, ii 872

Hermophenyl—A white amorphous solid. Introduced as a disinfectant.—*P.J.* '01, ii 245

HYDRARGYRI CYANIDUM $\text{Hg}(\text{CN})_2$, eq 250.5—Colourless or white prismatic crystals.

It contains theoretically 79.36 p c of metallic Mercury. It should be kept in well-stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light.

Solubility.—1 in 18 of Water, 1 in 20 of Alcohol (90 p c)

Medicinal Properties—A powerful antiseptic. Used as a local application (5 to 15 grains in 1 fl oz of Water = 0.3 to 1 gramme in 28.4 cc) to syphilitic rashes and sores of the throat, tongue, etc.—*Ringer*

In syphilis.—*P.J.* '95, ii 91 $\frac{1}{2}$ p c solution as an antiseptic.—*P.J.* '96, ii 19.

Subconjunctival and intravenous injections, in the treatment of serous syphilitic disease of the eye.—*B.M.J.* '03, ii 269

A lotion containing 0.25 gramme per 1000 grammes of Water used in acute conjunctivitis.—*M.P.* '05, ii 863

Solutions of the Cyanide or Oxycyanide are used for intravenous injection (*M.P.* '06, i 149), as they do not coagulate the blood. One cc of a solution containing Mercury Cyanide, 1 gramme, Distilled Water, 100 grammes, is injected daily in syphilis. The intravenous injections appear to act very promptly, but the most absolute sepsis must be insisted on.

Dose—Internally $\frac{1}{16}$ to $\frac{1}{8}$ grain = 0.004 to 0.008 gramme

Ph. Ger. maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme

Foreign Pharmacopœias—Official in Belg (*Cyanuretum Hydrargyri*), Fr (*Cyanure Mercurique*), Ger, Hung and Russ (*Hydrargyrum Cyanide*), Port (*Cyaneto Mercurico*), Mex. (*Cianuro de Mercurio*)

Tests—Mercuric Cyanide is decomposed on heating into metallic Mercury and Cyanogen gas, which burns with a purple flame. The aqueous solution is neutral in reaction towards Litmus paper. On the addition of Hydrochloric Acid it evolves the characteristic and highly poisonous odour of Hydrocyanic Acid. Neither Potassium Hydroxide Solution nor Ammonium Sulfate Potassium Iodide Solution yields a precipitate. On the addition of Hydrochloric Acid, when the solution behaves in a similar manner to Mercuric Chloride Solution, Hydrocyanic Acid Solution yields a black precipitate soluble in Ammonium Hydroxide Solution and in diluted Nitric Acid, Stannous Chloride Solution yields at first a whitish precipitate of Mercurous salt and subsequently a grey deposit of metallic Mercury. When gently heated with an equal part of Iodine in a dry test-tube it yields in the lower portion of the tube a yellow sublimate subsequently becoming red, and in the upper portion a colourless needle-shaped crystalline deposit.

The more generally occurring impurities are Mercuric Chloride and mineral residue. A delicate test for the former is to add to the 5 p c aqueous solution faintly acidified with Nitric Acid, one or two drops of Silver Nitrate Solution, no precipitate or turbidity should result. A solution of similar strength to the above

Pilula Hydrargyri Carbolicæ—Mercury Carbolate, $\frac{1}{2}$ grain, Extract of Liquorice, 1 grain, Powdered Liquorice, 1 grain, in each pill

Dose—Two to four pills daily

Hydrargyrol (Mercury Phenol para sulphonate)—Brownish red crystalline scales or crusts Decomposed by Water with the formation of basic salts Insoluble in Alcohol (90 p c) Introduced as an antiseptic

A combination of the above salt with Ammonium Tartrate is known under the name of 'Asterol,' a white or reddish white micro crystalline powder, soluble in Water Introduced as an antiseptic, used in the form of 2 to 5 p c solution—*B M J E* '01, 11 64, *P J* '99, 1 588, '99, 11 216, *C D* '01, 11 872

Hermophenyl (Sodium Mercuro-phenol Disulphonate)—A white amorphous powder, readily soluble in Water It contains about 40 p c Mercury Introduced as an antiseptic—*P J* '01, 11 245

HYDRARGYRI CYANIDUM $\text{Hg}(\text{CN})_2$, eq 250.5—Colourless or white prismatic crystals

It contains theoretically 79.86 p c of metallic Mercury It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light

Solubility—1 in 13 of Water, 1 in 20 of Alcohol (90 p c)

Medicinal Properties—A powerful antiseptic Used as a local application (5 to 15 grains in 1 fl oz of Water = 0.8 to 1 gramme in 28.4 cc) to syphilitic rashes and sores of the throat, tongue, etc—*Ringer*

Intravenous injection in syphilis—*P J* '95, 11 91 $\frac{1}{2}$ p c solution as an antiseptic in ophthalmic practice—*P J* '96, 11 19

Subconjunctival and intravenous injections, in the treatment of serous syphilitic disease of the eye—*B M J* '03, 11 269

A lotion containing 0.25 gramme per 1000 grammes of Water used in acute conjunctivitis (*M P* '05, 11 303)

Solutions of the Cyanide and Oxycyanide are used for intravenous injection (*M P* '06, 1 149), as they do not coagulate the blood One cc of a solution containing Mercury Cyanide, 1 gramme, Distilled Water, 100 grammes, is injected daily in syphilis The intravenous injections appear to act very promptly, but the most absolute sepsis must be insisted on

Dose—Internally $\frac{1}{16}$ to $\frac{1}{8}$ grain = 0.004 to 0.008 gramme

Ph Ger maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme

Foreign Pharmacopœias—Official in Belg (Cyanuretum Hydrargyri), Fr (Cyanure Mercurique), Ger, Hung and Russ (Hydrargyrum Cyanæ), Port (Cyaneto Mercurico), Mex (Cianuro de Mercurio)

Tests—Mercuric Cyanide is decomposed on heating into metallic Mercury and Cyanogen gas, which burns with a purple flame The aqueous solution is neutral in reaction towards Litmus paper, on the addition of Hydrochloric Acid it evolves the characteristic and highly poisonous odour of Hydrogen Cyanide, neither Potassium Hydroxide Solution nor Ammonia Solution yields a precipitate, Potassium Iodide Solution yields no precipitate until after the addition of Hydrochloric Acid, when the solution behaves in a similar manner to Mercuric Chloride Solution, Hydrogen Sulphide Solution yields a black precipitate, insoluble in Ammonium Hydrosulphide Solution and in diluted Nitric Acid, Stannous Chloride Solution yields at first a whitish precipitate of Mercurous salt and subsequently a grey deposit of metallic Mercury When gently heated with an equal part of Iodine in a dry test tube it yields in the lower portion of the tube a yellow sublimate subsequently becoming red, and in the upper portion a colourless needle shaped crystalline deposit

The more generally occurring impurities are Mercuric Chloride and mineral residue A delicate test for the former is to add to the 5 p c aqueous solution faintly acidified with Nitric Acid, one or two drops of Silver Nitrate Solution, no precipitate or turbidity should result A solution of similar strength to the above

should yield no reddish precipitate soluble in an excess of the reagent on the gradual addition of Potassium Iodide solution. Mineral residue is indicated by the ash left when the sample is ignited with free access of air.

Injectio Hydrargyri Cyanidi (Intravenous) —Mercuric Cyanide, 1 p c , inject 20 minims. —*Lot*

Mercury Oxycyanide as an antiseptic, in aqueous solution, 1 in 200 — *B M J E* '95, ii 104, *T G* '96, 405

Mercury Oxycyanide is official in Mex

MERCURY ZINCO-CYANIDE —A product which has been found by Lord Lister to have valuable anti-septic properties. —*P T* (3) xx 653, (3) xxii 769

There is also a gauze prepared with it —*B M J* '89, ii 1025, *L* '89, ii 943
Mercurialism resulting from use of the Cyanide gauze as a dressing —*P J* '96, ii 382

Unguentum Hydrargyri et Zinci Cyanidi —Mercury Zinc Cyanide, 2, 4 or 8 grains, Soft Paraffin, 1 oz —*London Ophthalmic*

HYDRARGYRI ETHYLENEDIAMINE CITRAS (Mercuramine) —A clear, colourless liquid, stated to be a 10 p c aqueous solution of Mercury Citrate containing 4 p c Ethylenediamine

Introduced as an antiseptic —*B M J* '01, ii 85, *P J* '01, ii 142

Under the title of **Sublamin** a combination of Mercury Sulphate and Ethylenediamine has been introduced. A 8 p c solution has been recommended as a disinfectant for the hands —*B M J E* '02, i 56

This salt is stated (*B M J* '05, i 727) not to injure the skin or discolour steel instruments in alcoholic solution. As a 2 in 100 alcoholic solution it gave results in sterilisation of the hand superior to any claimed for other methods, especially with regard to the power of penetration. May be conveniently kept as a 10 p c solution in Alcohol (50 p c)

HYDRARGYRI GALLAS (Mercury Gallate) —A dark grey or greyish-green amorphous powder, insoluble in Water. Is stated to be a more effective antiseptic than the Tannate. Used in syphilis

Dose. — $\frac{1}{2}$ to 1 grain = 0.016 to 0.06 gr., in a pill

HYDRARGYRI NAPHTHOLACETAS —Colourless, needle-shaped crystals, or as a white amorphous powder, insoluble in Water, has been used in syphilis

Dose — $\frac{1}{2}$ to 1 grain = 0.032 to 0.06 gr., in a pill

HYDRARGYRI SALICYLAS ($\text{HgCl}_2 \cdot \text{H}_2\text{O}$, eq 333.81) —A white or whitish amorphous odourless powder, practically insoluble in Water, and in Alcohol (90 p c). It is insoluble in 5 p c of metallic Mercury. It is soluble in Potassium or Sodium Hydroxide solution with the production of double salts. It is not dissolved in the cold by halogen compounds of the alkali metals, but dissolves on warming and when the solution cools double salts crystallise out.

Employed internally and by hypodermic injection, also as a dusting powder, in syphilis. Is stated to be as powerful a skin antiseptic as corrosive sublimate. An injection of 0.05 grammes recommended —*B M J* '04, ii 69 in tables

Intramuscular injection or Mercury Salicylate, 5, Liquid Paraffin, 50, one Pravaz syringe-full in syphilis —*B M J* '09, i 122

Maximum single dose, 0.02 grammes maximum daily dose, 0.06 grammes

In a large number of cases of syphilis intramuscular injections of 1 c c of the following emulsion Mercury Salicylate M 1, Liquid Paraffin, 10. Injections once a week, and not more than six injections, followed by an interval of rest of about a couple of months —*B M J* '04, i 609, ii 816

Foreign Pharmacopœias —Official in Ger, Jap, Mex, Russ, Swed and Swiss

Tests —Mercuric Salicylate does not answer the tests distinctive of Mercuric salts. It yields no precipitate with Hydrogen Sulphide or Ammonium Hydro-sulphide. It is decomposed by concentrated Hydrochloric, Nitric, and Sulphuric Acids, the solutions then yielding the distinctive tests given under Mercury. It yields a sublimate of metallic Mercury when heated in a dry

test tube A saturated aqueous solution yields with Ferric Chloride T S a violet coloration The method adopted by the *PG* for the determination of the Mercury is to mix a weighed quantity of 0.3 gramme with ten times its weight of Sodium Chloride and to dissolve the mixture in 100 c.c. of boiling Water, diluting the resulting solution to 400 c.c. The solution when slightly acidified with Hydrochloric Acid shall yield when completely precipitated with Hydrogen Sulphide 0.2 gramme of Mercuric Sulphide corresponding to 57.4 p.c. of metallic Mercury and 96.4 p.c. of pure Mercuric Salicylate An alternative method is to dissolve the salt in 8.5 c.c. of Nitric Acid and 13 c.c. of Hydrochloric Acid, evaporate to dryness, the residue is rendered acid with Hydrochloric Acid dissolved in Water, filtered, and the filtrate precipitated with Hydrogen Sulphide

The more generally occurring impurities are free Salicylic Acid and Sodium Salicylate The former may be detected by the marked acid reaction of the salt towards a piece of moistened blue Litmus paper, the latter by any residue remaining when the specimen is ignited with free access of air

HYDRARGYRI SUCCINIMIDUM ($\text{Hg}(\text{C}_4\text{H}_4\text{O}_2\text{N})_2$, eq 398.48) — White crystalline powder, soluble in Water It should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from the light It contains theoretically 50.5 p.c. w/w. of metallic Mercury Its solutions are stated not to precipitate albumen, and are therefore useful for hypodermic use Used as a solution of Mercury Succinimide, 38.2 grains, Cocaine Hydrochloride, 15.1 grains, Distilled Water, 775 grains — *L* '02, 1712

The use of Cocaine Nitrate in place of the Hydrochloride would avoid the precipitation of Calomel

Dose — $\frac{1}{2}$ to $\frac{1}{4}$ grain = 0.008 to 0.016 gramme

Foreign Pharmacopœias — Official in Ital

Tests — Mercuric Succinimide yields a solution which gives with Potassium Iodide Solution a bright scarlet precipitate soluble in excess of the reagent, when acidified with diluted Hydrochloric Acid, Hydrogen Sulphide yields a black precipitate, insoluble in Ammonium Hydrosulphide Solution, and in hot diluted Nitric Acid Solution, a piece of bright Copper foil immersed in an acidified solution of the salt becomes coated with a bright metallic film, with Stannous Chloride Solution a grayish-white precipitate is produced changing to gray, with Albumen solution no precipitate is produced When ignited with free access of air no weighable residue should remain

HYDRARGYRI SULPHAS (Mercuric Sulphate) *Syn* — **HYDRARGYRI PRÆSULPHAS**, SULPHATE OF MERCURY

A white, heavy, crystalline powder, HgSO_4 , eq 294.14, prepared by dissolving Mercury in strong Sulphuric Acid and evaporating to complete dryness It contains theoretically 67.6 p.c. w/w. of metallic Mercury It is decomposed by Water, forming a yellow oxysulphate called Turpeth Mineral ($\text{HgSO}_4 \cdot 2\text{H}_2\text{O}$), and free Sulphuric Acid

Foreign Pharmacopœias — Official in Fr (Sulfate Mercurique Basique), Mex Port and Span (Sulfato Mercurico) Not in the others

Unguentum Hydrargyri Sulphatis Flavæ (Turpeth Mineral Ointment Bazin's Ointment) — Yellow Mercury Sulphate, 15 grains, Benzoated Lard, 1 oz

Useful in ringworm and seborrhœa capitis

HYDRARGYRI TANNAS — A greyish green or blackish grey powder, containing 40 to 50 p.c. of Mercury

It should be preserved in well-closed bottles of a dark amber tint and protected as far as possible from the light

It is decomposed by Water and the solutions of the alkalis It is not materially affected by Diluted Hydrochloric Acid

Medicinal Properties — Very useful in syphilis

It is decomposed by the alkali of the intestines, and the Mercury rapidly passes into the system — *L* '84, 1723, *M T* '85, 1869

Dose—1 to 2 grains = 0.06 to 0.13 gram, in a pill, three times a day, an hour before meals

Foreign Pharmacopœias—Official in Aust. (Hydrargyrum tannicum oxydulatum), contains about 55 to 57 per cent of Mercury, Mex (Tanato de Mercurio) Not in the others

Tests—Mercury Tannate is not materially affected by dilute Hydrochloric Acid, but the concentrated acid decomposes it with formation of Mercurous Chloride and Tannic Acid. It is decomposed by alkali Hydroxide solutions and solutions of alkali carbonates, the alkaline solution rapidly darkening on exposure to air. It should be free from Nitrates as indicated by rubbing 0.8 gramme of the salt with 3 c.c. of Water, filtering and adding two drops of the filtrate to 5 c.c. of Diphenylamin. When no blue colour should be produced. When ignited with free carbon leaves no weighable residue.

HYDRARGYRI THYMOLACETAS—A white micro-crystalline powder, almost insoluble in Water. Has been used as an intra-muscular injection (10 per cent in liquid paraffin).

Dose— $\frac{1}{2}$ to 1 grain = 0.032 to 0.06 gram

HYDRARGYRI IODIDUM RUBRUM.

MERCURIC IODIDE

B.P. Syn.—BINIODIDE OF MERCURY

HgI₂, eq. 450.60

FR, BI-iodure de Mercure, GER, Jodkalium, ITAL, BIODURO DI MERCURIO, SPAN, Yodo de Mercurio

Scarlet-red crystals, or a scarlet-crystalline powder

It should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from light.

It contains theoretically 44.1 per cent of Mercury.

Solubility.—Almost insoluble in Water, sparingly soluble in Glycerin, 1 in 300 of Alcohol (90 per cent), 1 in 70 of Ether, 1 in 280 of Olive or Almond Oil or Lard, 1 in 50 of Castor Oil. Freely in an aqueous solution of Potassium Iodide or Mercuric Chloride.

Medicinal Properties.—Also a powerful irritant poison in over-doses, similar to the Green Iodide, only much more active. It is used internally in the same cases as Corrosive Sublimite, more particularly in chronic glandular enlargements and rheumatism and cutaneous diseases when due to syphilis. As an antiseptic lotion (1 in 5000) in surgical and obstetric practice.

The Ointment is a most effective application for bronchocœle, and a good application for warts and syphilitic nodes and for lupus. If applied to the eyelids, should be diluted to quarter the strength.

In infantile diarrhoea—Pr. 1920, P.J. '95, 11 215

Has been used (L. '04, 11 1396) in the form of a 1 in 1000 Chloroform solution for the sterilisation of cat-gut ligatures.

Recommended (B.M.J. '05, 11 785) for the sterilisation of the hands before surgical operation, a 1 in 500 solution obtained for two minutes by means of gauze, the hands being previously washed in very hot water and dehydrated by means of alcohol.

Dose.— $\frac{1}{32}$ to $\frac{1}{16}$ grain = 0.002 to 0.004 gramme

Ph Ger maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme

Prescribing Notes—*Usually given in the form of Pilules well triturated with Milk Sugar and 'Diluted Glucose' When prescribed in Solution it is dissolved by the aid of Potassium Iodide It can also be dissolved in Castor Oil and given in Capsules*

Official Preparation—Unguentum Hydrargyri Iodidi Rubri Used in the preparation of Liquor Arseni et Hydrargyri Iodidi

Not Official—Hydrargyri et Potassii Iodidum, Injectio Hydrargyri Iodidi Rubri and Unguentum Hydrargyri et Potassii Iodidi (Lutz's Ointment)

Foreign Pharmacopœias—Official in Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap. (Hydrargyrium Biniodatum), Mex., Port., Russ., Span., Swiss and U.S.

Tests—Mercuric Iodide becomes yellow when heated, but again assumes its scarlet colour on cooling, the *U.S.P.* specifies the temperature 150° C (302° F). When heated with Potassium Hydroxide Solution and a little Milk Sugar it yields a grayish precipitate of metallic Mercury. If this precipitate be well washed and dissolved in a mixture of Nitric and Hydrochloric Acid, it yields on neutralisation of the excess of acid the tests distinctive of Mercuric salts given under Hydrargyri Perchloridum. A portion of the filtrate, when slightly acidified with diluted Nitric Acid, yields with Silver Nitrate Solution a curdy yellow precipitate, insoluble in Nitric Acid, almost insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution. Another portion of the filtrate acidified with Hydrochloric Acid affords, on the addition of Chlorine Water, or Sodium Nitrite Solution, a reddish-yellow colour, soluble to a violet coloured solution in Carbon Bisulphide. The *B.P.* states that when heated with excess of Copper it should yield from 43.5 to 44 p.c. of metallic Mercury, but gives no indication as to whether the Mercury is to be gravimetrically or volumetrically determined. The *U.S.P.* states that it should contain not less than 98.5 p.c. of pure Mercuric Iodide, but gives no method of determination.

The more generally occurring impurities are Mercurous Iodide, Mercuric Chloride, soluble Chlorides or Iodides, and mineral matter. The absence of Mercurous Iodide may be ensured by the ready and complete solubility of the specimen in Ether, and in Potassium Iodide Solution, Mercuric Chloride, by Alcohol (94.9 p.c.) and Litmus Test described below, soluble Chlorides and Iodides, by the Hydrogen Sulphide and the Silver Nitrate Tests also given. When ignited with free access of air the salt leaves no weighable residue.

Alcohol and Litmus—The cooled alcoholic solution of the salt should be colourless, and should not redden blue Litmus paper, *P.G.*, a saturated solution of the salt in hot Alcohol (94.9 p.c.), when cooled and diluted with an equal volume of Water should not redden blue Litmus paper, *U.S.P.*

Hydrogen Sulphide—If the salt be thoroughly agitated with Water (0.5 gramme with 10 c.c., *U.S.P.*) and filtered, the filtrate should be only slightly coloured by T.S. of Hydrogen Sulphide, *P.G.* and *U.S.P.*

Silver Nitrate—The filtrate obtained as above should only be rendered slightly opalescent with T.S. of Silver Nitrate, *P.G.* and *U.S.P.*

Preparations.**UNGUENTUM HYDRARGYRI IODIDI RUBRI MERCURIC IODIDE OINTMENT** *B P Syn*—OINTMENT OF RED IODIDE OF MERCURY

Mix 1 of Mercuric Iodide, in fine powder, with 24 of Benzoated Lard (1 in 25)

Foreign Pharmacopœias—Official in Fr, Mercuric Iodide 1, Lard 8, Mex., Pomada, 1 in 50 Not in the others

Not Official**HYDRARGYRI ET POTASSII IODIDUM**—Yellow acicular crystals

It is a powerful antiseptic

INJECTIO HYDRARGYRI BINIODIDI (pro Vagina)—Mercuric Chloride 8 grains, Potassium Iodide 5 grains, Water to 1 fl oz, 1 fl dr to a pint of Water = 1 in 10,000—*Lock*

It requires 22 grains of the Potassium Iodide to form a solution

INJECTIO HYDRARGYRI IODIDI RUBRI—Red Iodide of Mercury, 5 grains, Iodide of Potassium, 20 grains, Water to 20 fl oz—*St Bartholomew's* (1 in 1750)

Mercuric Iodide, 1, Potassium Iodide, 4, Distilled Water, q s to produce 100 The dilution of this solution to 100 times its volume forms a 1 in 10,000 solution of Mercuric Iodide—*B P C*

UNGUENTUM HYDRARGYRI ET POTASSII IODIDI (Lutz's Ointment)—Red Mercuric Iodide, 5 grains, Potassium Iodide, 5 grains, Water, q s, Prepared Lard, 1 oz

University has a similar preparation containing Wool Fat

Not Official.**HYDRARGYRI IODIDUM VIRIDE.**

GREEN IODIDE OF MERCURY GREEN MERCUROUS IODIDE

HgI, eq 324 73

A dull green powder containing excess of Mercury, which decomposes upon exposure to light

It should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from the light

It has been shown (*P J '00*, 11 87) that the intensity of the green colour naturally depends upon the relative proportions or excess of Mercury employed. The yellow Mercurous Iodide of the *U S P* is shown to be quite uniform in composition and also its colour is stable when properly protected. It is naturally for the therapist to decide whether a Mercurous Iodide containing more or less free Mercury is preferable to the pure salt for medicinal use—*C D '00*, 11 164, *P J '00*, 11 86

Solubility—Insoluble in Water, Alcohol, and Ether

Medicinal Properties—Given in syphilis and in tubercular and rheumatic affections. Employed as an ointment (1 part to 8 of Lard) for syphilitic eruptions, chronic skin diseases, enlarged glands, and bronchocœle

Dose—It varies with different prescribers from $\frac{1}{2}$ grain to 2 grains = 0.01 to 0.13 gramme

Ital maximum single dose, 0.05 gramme, maximum daily dose, 0.20 gramme

$\frac{1}{2}$ or $\frac{1}{4}$ grain three times daily, increasing until constitutional effects are produced—*L '01*, 11. 1038.

Prescribing Notes — *It makes a good pill with Sugar of Milk and 'Diluted Glucose'*

Incompatible with soluble Iodides — *C D '92, 11 275*

Foreign Pharmacopœias — Official in Austr, Hung, Swiss and US, (*Hydrargyrum Iodatum flavum*), Belg (*Proto Ioduretum Hydrargyri*), Dutch and Swed (*Iodetum Hydrargyrosus*), Fr (*Proto-Iodure de Mercure*), Ital (*Ioduro Mercurioso*), Jap, (*Hydrargyrum Iodatum*), Mex (*Ioduro Mercurioso*), Port (*Iodeto Mercurioso*), Swiss (*Hydrargyrum Iodatum*), Span (*Ioduro Mercurioso*)

Tests — Mercurous Iodide melts when heated, and is entirely volatilised at a red heat. Dissolved in Nitric Acid it yields a solution answering the tests characteristic of Mercury given under *Hydrargyrum*. Under the influence of light it undergoes decomposition, with formation of Mercuric Iodide and metallic Mercury. Heated with Manganese Dioxide and Sulphuric Acid it evolves violet vapours of Iodine. When shaken in a dry test tube with purified Ether, filtered, and the Ether evaporated, no residue should remain indicating the absence of Mercuric Iodide.

The yellow Mercurous Iodide is official in the *USP* and is required to contain not less than 99.5 p.c. of pure Mercurous Iodide. The absence of more than traces of Mercuric Iodide is ensured by shaking 0.5 gramme of the salt with 10 cc of Alcohol (94.9 p.c.) allowing to stand, and filtering, portions of the perfectly clear filtrate should be scarcely affected by Hydrogen Sulphide, should yield only a faint opalescence when dropped into Water, and only a faint red stain when evaporated in a white porcelain dish.

PILULA HYDRARGYRI IODIDI VIRIDIS — Green Mercurous Iodide, $\frac{1}{2}$ grain, Opium, $\frac{1}{2}$ grain, Extract of Gentian, 2 grains

PILULE D'IODURE MERCUREUX OPIACÉES *Pilules de Record (Fr)* — Recently prepared — Mercurous Iodide, 0.5 gramme, Powdered Opium, 0.2 gramme, Liquorice Powder, 0.3 gramme, Honey *q s*, divide into 10 pills

UNGUENTUM HYDRARGYRI IODIDI VIRIDIS CUM ATROPINA — Green Mercurous Iodide, 10 grains, Atropine, 1 grain, Lard, $\frac{1}{2}$ oz

HYDRARGYRI NITRATIS LIQUOR ACIDUS.

ACID SOLUTION OF MERCURIC NITRATE

A heavy, colourless, strongly acid solution, containing about 33 p.c. of Mercury in the form of Mercuric Nitrate. It is obtained by dissolving, in the cold, 4 (by weight) of Mercury in 5 of Nitric Acid diluted with $1\frac{1}{2}$ of Water. It should be preserved in well-stoppered amber-tinted glass bottles.

Medicinal Properties — Caustic and antiseptic. Applied to syphilitic warts, ulcers, etc., care should be taken that the surrounding healthy parts are not touched. Used in cancerous growths and in lupus. As a gargle, 1 or 2 minims to 1 fl oz Water. As an injection in gonorrhœa, 1 minim to 2 fl oz Water.

Official Preparations — *Unguentum Hydrargyri Nitratis* and *Unguentum Hydrargyri Nitratis Dilutum* contain Mercuric Nitrate.

Foreign Pharmacopœias — Official in US (*Liquor Hydrargyri Nitratis*) prepared from Mercuric Oxide, sp. gr. about 2.086 at 25° C (77° F), Fr (*Azotate de Bioxyde de Mercure Dissous*), sp. gr. 2.246, Ital (*Nitrato Mercurico liquido*), sp. gr. 2.250, Port (*Soluto de Azotato Mercurico*), Span (*Nitrato Mercurico Acido*), sp. gr. 2.246, Mex. (*Nitrato Mercurico*). Not in the others.

Tests.—Acid Solution of Mercuric Nitrate a sp gr of about 2.0. It yields when diluted the tests distinctive of Mercuric salts given under Hydrargyri Perchloridum. Ferrous Sulphate Solution carefully poured on to the surface of the solution yields a dark brown ring at the point of contact of the two fluids. It should be free from Mercurous salts as ascertained by the non-appearance of a precipitate or cloudiness, when the solution is diluted with Water, or on the addition of diluted Hydrochloric Acid. When evaporated to dryness and ignited with free access of air no weighable residue should remain.

UNGUENTUM HYDRARGYRI NITRATIS. MERCURIC NITRATE OINTMENT *B.P. Syn*—OINTMENT OF NITRATE OF MERCURY *N.O. Syn*—CITRINE OINTMENT

Mercury (by weight), 1, Nitric Acid, 3, Lard, 4, Olive Oil (by weight), 7. Dissolve the Mercury in the Nitric Acid without the aid of heat, agitating gently from time to time. Heat the Lard and Olive Oil together on a sand-bath, so that the mixture when transferred to a heated earthenware jar, capable of holding 10 times the quantity, shall be at a temperature of about 290° F (143.3° C). Add the cold Mercurial Solution very gradually, stirring constantly to promote disengagement of the fumes. After frothing has ceased, the mixture, which should have a temperature of not less than 200° F (93.3° C), must be kept stirred until it is cold. The resulting Ointment should be firm in consistence and have a pale lemon colour.

(about 1 in 16½)

The official directions given above do not work satisfactorily, the temperature is much too high and yields varying results with different operators, and even by the same operator at different times. The following method will yield a more uniform product.

Dissolve the Mercury in the Nitric Acid without the aid of heat. Heat the Lard and Oil on a water-bath, until the Lard is dissolved and at a temperature of 52.2° to 87.8° C (120° to 190° F) add the Mercuric Solution (cold) to the melted fats and stir continuously. When brisk effervescence has commenced continue the heat for 10 minutes, then remove from the water-bath and stir till cold.

The product should have a good consistence, and if kept in covered pots should retain its pale lemon colour for several months. In the hands of the author this method has never yielded a 'spongy' product. The heat should not be continued until all action has ceased for the product will then be of a darker colour and blacken in the course of a week or two—*P.J.* 97, 1 172, '98, 11 165, 179, 232, 236, *C.D.* '93, 1 933, *A.J.P.* 97, 203, 232.

Two specimens made by the above process in 1898 were exhibited at an evening meeting in 1902 on account of their good condition of preservation. It is noted (*P.J.* '02, 1 314) that the two chief objections which have been raised to the Squire process are (a) that the Ointment must be, from the nature of the case, extremely acid, and (b) that however well stirred it is apt to be spongy. With regard to the former, it is shown that an Ointment prepared by the *B.P.* process contained the equivalent of 5.04 p.c. of Nitric Acid, whilst one prepared by 'Squire's process' indicated only 4.4 p.c., with regard to the latter, the percentage of spongy batches was greater in the case of the Ointment made by the official than in that made by 'Squire's process'.

It is contended (*P.J.* '02, 1 350) that the spongy character of these specimens were due to their initial excessive acidity, and that an Ointment should not necessarily be expected to retain its colour for several months.

period. Attention is again drawn to the fact (*P J* '02, 1 368) that an Ointment prepared by the *B P* 1898 process was more acid than one prepared by the Squire process, and that, with one or two exceptions, it is generally admitted that the latter gives more uniform results than the former process. It is still regarded (*P J* '02, 1 394) as inconceivable that a carefully made *B P* Ointment could by any possibility contain more acid than one made by 'Squire's process', and an explanation is asked for a specimen assaying 2.1 p.c. of acid. For a refutation of the opinion respecting the comparative acidities the reader is referred (*P J* '02, 1 436) to the experiments recorded *P J* '02, 1 314. The Mercury has generally been assumed to exist in the state of Mercuric Nitrate, and hence, presumably, the name given to it in the *B P*. A percentage of 2.1 corresponds very closely with Mercurous Nitrate.

Ointments prepared by the official process and by that recommended by the author have been critically compared (*P J* '04, 11 736). When freshly prepared there was little difference in them, the official being slightly darker. After six weeks the official Ointment had become slightly spongy. After a further six weeks the official was distinctly darker. After nearly five months that made by the author's process was still pale yellow in colour, whilst the official was distinctly darker. Although there is little to choose between the two methods, yet the evidence is somewhat in favour of Squire's method. The experiments repeated with ingredients obtained from different sources showed, in each case, a slight advantage in favour of Squire's method.

Medicinal Properties—Applied in diseases of the skin as a parasiticide, in *tinea tarsi* it is diluted with 7 parts of Vaseline and applied by means of a camel's-hair pencil to the eyelids. Diluted with Glycerin and applied by a brush to the nostrils in *ozæna*.

This Ointment, when diluted with Lard, soon acquires a leaden colour, it changes less with Spermaceti Ointment, and least of all when diluted with Soft Paraffin.

Incompatibles—All reducing agents, Camphor, Essential Oils, Lard, etc.

Official Preparation—*Unguentum Hydrargyri Nitratis Dilutum*

Not Official—*Unguentum Metallorum*, *Unguentum Hydrargyri Zinci et Plumbi*

Foreign Pharmacopœias—Official in Belg., Mercury 5, Nitric Acid 7, Lard 45, Olive Oil 43, Fr., Mercury 1, Nitric Acid (sp. gr. 1.394) 2, Lard 10, Olive Oil 10, Mex., Mercury 4, Nitric Acid 6, Lard 64, Port., Sol. Mercuric Nitrate 2, Lard 9, Olive Oil 9, Swed., Mercury 1, Nitric Acid (sp. gr. 1.5) 2, Lard 12, U.S., Mercury 7, Nitric Acid (sp. gr. 1.414) 17.5, Lard 76.

UNGUENTUM HYDRARGYRI NITRATIS DILUTUM.
DILUTED MERCURIC NITRATE OINTMENT *B P Syn*—DILUTED OINTMENT OF NITRATE OF MERCURY

Mix 1 of Mercuric Nitrate Ointment with 4 of yellow Soft Paraffin.
(1 in 5)

It is more dilute than *B P* '85

Not Official.

UNGUENTUM METALLORUM—Mercuric Nitrate Ointment, Lead Acetate Ointment, Zinc Ointment, equal parts —*King's and Great Northern*

UNGUENTUM HYDRARGYRI ZINCI ET PLUMBI *Syn*—**UNGUENTUM METALLORUM**—Mercurous Chloride, 10 grains, Mercuric Nitrate Ointment, 20 grains, Lead Acetate, 10 grains, Zinc Oxide, 20 grains, Soft Paraffin (yellow), to 1 oz.—*St Thomas's*

This has been incorporated in the *B P C* as follows—

Mercurous Chloride, 2, Mercuric Nitrate Ointment, 4, Lead Acetate, in powder, 2, Zinc Oxide, finely sifted, 4, Soft Paraffin (yellow), sufficient to produce 100.

HYDRARGYRI OLEAS.**MERCURIC OLEATE**

FR, OLEATE DE MERCURE, GFR, OELSÄURES QUECKSILBER

A brownish-yellow semi-solid oleaginous mass when fresh, but becoming of a stiffer consistence and darker colour on keeping, it should therefore be kept in well-stoppered glass bottles of a dark amber tint and exposed as little as possible. It is the precipitate obtained on mixing sol. ion of Mercuric Chloride and Hard Soap

An Oleate containing 20 p.c. is readily made as follows — Mercuric Oxide (finely powdered), 4 Oleic Acid (by wt. 1 lb. 16, 1.740 (0.720), 1 Mix the Oxide of Mercury with the Ether and stir rapidly in a whole of the Oleic Acid, warm to 120° F, stirring frequently until the Oxide is dissolved. The operation should be complete in 1 to 2 hours

This method has been incorporated in the *B.P.C.* under the title *Oleinatum Hydrargyro*

A revelation to the method of direct combination of Mercuric Oxide and Oleic Acid has been recommended

Mercuric Oleate was introduced by Prof. Marshall in 1872, and was made of three different strengths, containing respectively 5 p.c., 10 p.c., and 20 p.c. of Mercuric Oxide

The 5 p.c. very quickly changed to a black colour, owing to reduction of the Mercuric Oxide, the 10 p.c. is not very long without change. It is better to keep the 20 p.c. and use as required for use

The Mercuric Oleate of the *U.S.P.* is prepared by the interaction of yellow Mercuric Oxide and Oleic Acid

Medicinal Properties — Similar to those of Mercury Ointment and Liniment, but more easily absorbed. Used with great success in tubercular peritonitis. Has been strongly recommended as an application for persistent inflammation in the joints or other parts near the surface, more particularly when combined with Morphine. It is useful, spread on lint and placed in the axilla, for syphilis, also as an application for non-ulcerative chancres. A good application for pityriasis versicolor.

Official Preparation — Unguentum Hydrargyri Oleatis

Not Official — Hydrargyri Oleas c. Morphina

Foreign Pharmacopœias — Official in Jap, Mex and U.S. Not in the others

Tests — Mercuric Oleate heated with a piece of bright Copper foil and a little dilute Hydrochloric Acid deposits a film of metallic Mercury. When dissolved in Ether shaken with diluted Nitric Acid, and the aqueous acid portion separated, the latter yields the tests distinctive of Mercuric salts given under Hydrargyri Perchloridi. The washed ethereal liquid transferred to a flask, the Ether removed by distillation, leaves a residue which consists mainly of Oleic Acid. A method for the determination of the Mercury is described *Y.B.P.* 1900, 200. A weighed quantity of 2 grammes of the Oleate is weighed into a flask, shaken with Ether and stirred with 10 cc of Ether until completely dissolved, then 25 cc of Alcohol (90 p.c.) and 5 cc of Hypophosphorous Acid are then added and the whole shaken. The residue is then reduced Mercury completely sublimed.

solution The liquid is poured off, the precipitate washed by decantation successively with Alcohol (90 p c) and Ether, the beaker and Mercury dried at 100° C (212° F), cooled and weighed The weight multiplied by 50 yields the percentage w/w of metallic Mercury The official Oleate was found to contain 23 p c The Oleate should leave no weighable residue upon ignition, when shaken with Water and filtered, the filtrate yields only a faint opalescence with Silver Nitrate Solution, and no marked darkening in colour with Hydrogen Sulphide

Preparation

UNGUENTUM HYDRARGYRI OLEATIS MERCURIC OLEATE OINTMENT

Mercuric Oleate, 1, Benzoated Lard, 3

Not Official

HYDRARGYRI OLEAS C MORPHINA is made by dissolving 1 grain of Morphine alkaloid in each drgm of Mercuric Oleate (10 p c)

HYDRARGYRI OXIDUM FLAVUM.

YELLOW MERCURIC OXIDE

HgO, eq 214 68

Fr, OXIDE DE MERCURE JAUNE, GER, GELBES QUECKSILBEROXYD, ITAL, OSSIDO GIALLO DI MERCURIO, SPAN, OXIDO MERCURIO AMARILLO

An orange-yellow heavy amorphous powder, being the precipitate obtained from solutions of Mercuric Chloride and Sodium Hydroxide It is important that it should be protected from light

Solubility—Practically insoluble in Water or Alcohol (90 p c) Asparagin dissolves the freshly precipitated Oxide (*see* p 125) to form **Mercury-Asparagin**

Medicinal Properties.—Similar to Red Mercuric Oxide

Ph Ger maximum single dose, 0 02 gramme, maximum daily dose, 0 06 gramme

Official Preparation—Unguentum Hydrargyri Oxidi Flavi

Foreign Pharmacopœias—Official in Austr, Hung, Jap and Swiss, (Hydrargyrum Oxydatum Flavum), Belg (Oxydum Hydrargyri Flavum), Dan and Dutch (Oxydum Hydrargyricum Flavum), Fr (Oxyde de Mercure Jaune), Ger and Russ (Hydrargyrum Oxydatum viâ Humidâ Paratum), Ital (Ossido Mercurico Giallo, Norw. (Oxidum Hydiargyricum Flavum), Mex and Span (Oxido Mercurico Amarillo), Swed (Oxydum Hydiargyricum Præcipitatum), US (Hyd Oxid Flav)

Tests—Yellow Mercuric Oxide when gently heated assumes a red colour, and at a dull red heat it is completely decomposed into metallic Mercury and Oxygen, the presence of the latter can be demonstrated by placing the glowing end of a match into the vessel in which it is being heated, the match immediately igniting The Oxide is readily and completely soluble in diluted Hydrochloric Acid,

yielding a solution which answers the tests distinctive of Mercuric salts given under Hydrargyri Perchloridi. The *BP* requires that the proportion of metallic Mercury obtained, presumably when heated to incipient redness, should be 92 to 92.5 p.c., corresponding to 99.3 to 99.9 p.c. of pure Yellow Mercuric Oxide, but does not state how the Mercury is to be collected or determined. The *USP* states that it should contain not less than 99.5 p.c. of pure Yellow Mercuric Oxide, but does not give a method of determination. The *PG* gives neither a percentage nor method of determination.

It may be distinguished from the Red Oxide by digestion for 15 minutes on a water-bath with twice its weight of Oxalic Acid dissolved in a small quantity of Water, when it will be converted into white Mercuric Oxalate. The *PG* test directs agitation with a 1 in 10 Oxalic Acid Solution.

The more generally occurring impurities are fixed residue, Chlorides, foreign salts, Arsenic, and foreign metals. The *BP* examines only for fixed residue. It is required to yield only an insignificant amount when heated to incipient redness, the *USP* states that at a red heat it is finally volatilised, leaving not more than 0.1 p.c. of residue, the *PG* that 0.2 gramme shall leave at most a residue which cannot be weighed. The solution in diluted Nitric Acid should yield only the slightest opalescence with Silver Nitrate Solution, indicating the absence of Chlorides. Foreign salts, metals, and Arsenic may be detected by the tests given under Hydrargyri Subchloridum.

Oxalic Acid—When it is agitated with Oxalic Acid Solution (1-10) it is gradually converted into a white crystalline powder, *PG*, when 0.5 gramme of Oxide and 1 gramme of Acid in 10 c.c. of Water is digested on a water-bath for 15 minutes the Oxide is converted into white Mercuric Oxalate (direction from Red Mercuric Oxide), *USP*.

Silver Nitrate—A solution in dilute Nitric Acid (1-50) should be clear and should not afford more than slight opalescence with *TS* of Silver Nitrate, *PG*, 0.1 gramme of Oxide dissolved in 10 c.c. of diluted Nitric Acid should not afford more than a slight opalescence with Silver Nitrate Solution, *USP*.

If 0.5 gramme of the Oxide be dissolved in a mixture of 2 c.c. of Hydrochloric Acid and 25 c.c. of Water, the solution should not respond to the *USP* tests for foreign salts, metals or Arsenic given under Hydrargyri Subchloridum, *USP*.

Preparation

UNGUENTUM HYDRARGYRI OXIDI FLAVI—*YELLO'W MERCURIC OXIDE OINTMENT*

Yellow Mercuric Oxide, in very fine powder, 10 grains, Yellow Soft Paraffin, 490 grains (1 in 50)

Medicinal Properties—Used in cases of chronic eczema, pityriasis, ringworm, chronic lichen, and syphilitic eruptions.

Diluted with an equal or twice the quantity of Vaseline, is a most valuable remedy for ophthalmia, tarsi, corneal ulceration, and all forms of conjunctival inflammation.

A 4 p.c. Ointment of the Yellow Oxide in keratitis profunda.

11 1308

Several formulas are given in the Pharmacopoeia for the preparation of ointments containing from 2 to 60 grains to the ounce.

Foreign Pharmacopœias.—Official in Belg, Yellow Oxide 1, Vaseline 49, Dutch, Yellow Oxide 1, White Vaseline 19, Fr (Pommade d'Oxyde de Mercure Jaune), Yellow Oxide 1, Vaseline 19, Mex (Pomada de Oxido Amarillo de Mercurio), Yellow Oxide 1, Vaseline 15, Ital (Pomata di ossido giallo di mercurio), Yellow Oxide 1, Vaseline 153, Jap, Yellow Oxide 1, Vaseline 9, Russ, Yellow Oxide 1, Lard 49, Span, Yellow Oxide 1, Vaseline 19, Swiss, Freshly precipitated Mercuric Oxide 5 in Water 15, Wool Fat 20, Vaseline 60, U S, Yellow Mercuric Oxide 1, Water 1, Hydrus Wool Fat 4, Petrolatum, 4

HYDRARGYRI OXIDUM RUBRUM.

RED MERCURIO OXIDE

HgO , eq 214 68

FR, OXYDE DE MERCURE ROUGE, GER, QUECKSILBEROXYD, ITAL, OSSIDO MERCURICO ROSSO, SPAN, OXIDO MERCURIO ROJO

Orange-red crystalline scales, or heavy crystalline orange-red powder, prepared from Mercurous Nitrate

It should be kept in well-closed glass bottles of an amber tint and protected as far as possible from the light

Solubility—Insoluble in Water and Alcohol (90 p c), readily soluble in Hydrochloric Acid

Medicinal Properties—A powerful irritant, rarely used internally Employed in form of ointment, *q v*

Ph Ger maximum single dose, 0 02 gramme, maximum daily dose, 0 06 gramme

Official Preparation—Unguentum Hydrargyri Oxidi Rubri

Foreign Pharmacopœias—Official in U S, Belg (Oxydum Hydrargyri Rubrum), Dan, Dutch and Norw (Oxydum Hydrargyricum), Fr (Oxyde de Mercure Rouge), Ger and Swiss (Hydrargyrum Oxydatum), Ital (Ossido Mercurico Rosso), Jap (Hydrargyrum Oxydatum Rubrum), Mex (Oxido Mercurico), Port (Oxyde Mercurico), Russ (Hydrargyrum Oxydatum Levigatum), Span (Oxido Mercurico Rojo) Not in Austr or Hung

Tests—Red Mercuric Oxide undergoes change of colour when heated, turning a dark violet or almost black, but regains its original orange-red colour on cooling The *U S P* specifies a temperature 400°C (752°F) At a red heat it is decomposed into metallic Mercury and Oxygen, the presence of the latter can be ascertained in the same way as described under Hydrargyri Oxidum Flavum It is readily and completely soluble in diluted Hydrochloric Acid, yielding a faintly opalescent solution which answers the tests distinctive of Mercuric salts given under Hydrargyri Perchloridum It is officially required to answer the tests given in the *B P* under Hydrargyrum Oxidum Flavum, consequently the same remarks as are there made respecting the requisite official percentage of Mercury apply also here The *U S P* requires it to contain not less than 99 5 p c of pure Red Mercuric Oxide, but gives no method of determination The *P G* gives neither a percentage nor a method of determination It may be distinguished from the Yellow Oxide by remaining unaltered when

treated with Oxalic Acid Solution. The *U.S.P.* digests the Oxide with Oxalic Acid and a small quantity of Water on a water-bath, requiring that it shall undergo no change in colour in 2 hours. The *P.G.* repeatedly agitates the Oxide with a 10 p.c. w/w Oxalic Acid Solution, when it should not undergo material change in 15 minutes. The *B.P.* does not include a similar test. The respective tests will be found in detail in small type below.

The more generally occurring impurities are fixed residue, Chlorides, Nitrates, metals, foreign salts, or Arsenic. The *B.P.* requires that it shall leave only an insignificant amount of fixed residue when heated to incipient redness, the *U.S.P.* that it leaves no appreciable residue at a red heat, the *P.G.* that 0.2 gramme shall not leave a residue which cannot be weighed. Its solution in diluted Nitric Acid should yield only a slight opalescence at the most, with Silver Nitrate Solution. No test for Chlorides is included in the *B.P.* The *B.P.* test for Nitrates requires that it should not evolve orange fumes when heated in a dry test-tube; the *U.S.P.* and *P.G.* employ the Ferrous Sulphate and Sulphuric Acid Test. Metals, foreign salts, or Arsenic may be detected as described under Hydrargyri Subchloridum.

Oxalic Acid.—When Red Mercuric Oxide is repeatedly agitated with Oxalic Acid Solution (1-10) it should not undergo any material change in colour in 15 minutes, *P.G.*, 0.5 gramme of Oxide and 1 gramme of Oxalic Acid in 10 cc of Water, digested on a water-bath, the Mercuric Oxide should not change in colour in 2 hours, *U.S.P.*

Sulphuric Acid and Ferrous Sulphate.—I. 1 gramme of Red Mercuric Oxide be mixed with 2 cc of Water and 2 cc of Sulphuric Acid added, on the further addition of 1 cc of Ferrous Sulphate T.S. poured carefully upon it, there should be no coloured zone at the junction of the liquids even after standing for some time, *P.G.*, 1 gramme mixed with 5 cc of Water and 2 cc of Sulphuric Acid, cooled, and 2 cc of Ferrous Sulphate T.S. carefully poured on it no brown-coloured zone should be seen at the line of contact on standing, *U.S.P.*

Silver Nitrate.—The solution of the Oxide (1-50) obtained by means of a dilute Nitric Acid should be clear and should only become slightly turbid with T.S. of Silver Nitrate, *P.G.*, 0.1 gramme of the Oxide of diluted Nitric Acid should not produce more than a slight opalescence with T.S. of Silver Nitrate, *U.S.P.*

Preparations

UNGUENTUM HYDRARGYRI OXIDI RUBRI. RED MERCURIC OXIDE OINTMENT *B.P.*, *U.S.P.*, *P.G.*, PRECIPITATE OINTMENT

Red Mercuric Oxide, in very fine powder, $\frac{1}{2}$, Yellow Paraffin Ointment, 24 (1 in 10)

Medicinal Properties.—Stimulant for chronic ulcers and caustic for unhealthy granulations and soft warts, skin parasiticide. Much diluted, is used for ulcerations of the cornea and chronic ophthalmia, but the Ointment of the Yellow Oxide is preferred by many.

Foreign Pharmacopœias.—Official in Belg., 1 in 50, Dan., Dutch, Fr., Norw., Port. and Swiss, 1 in 20, Mex., Ger. and Ital., 1 in 16; Jap., Span. and U.S., 1 in 10; Russ., with Yellow Oxide (p. 519).

HYDRARGYRI PERCHLORIDUM.

MERCURIUM CHLORIDE

HgCl₂, eq 269 18FR, BICHLORURE DE MERCURE, GER, QUECKSILBERCHLORID, ITAL,
BICHLORURO DI MERCURIO, SPAN, CLORURO MERCURIO*B P Syn* —BICHLORIDE OF MERCURY, CORROSIVE SUBLIMATE, PERCHLORIDE OF MERCURY*N O Syn* —Chloretum Hydrargyricum, Hydrargyrum Bichloratum, Sublimatus Corrosivus

Heavy, colourless, rhombic crystals, or in crystalline masses, or a heavy white crystalline powder

Odourless, and possessing a particularly acrid and persistent metallic taste but should only be tasted with extreme caution.

Solubility —1 in 19 of Water, 1 in 5 of Alcohol (90 p c), 1 in 3 of Absolute Alcohol, 1 in 6 of Ether, *B P* (0 735), 1 in 11 of Purified Ether (0 720), 8 in 13 of Glycerin**Medicinal Properties** —A powerful antiseptic and very poisonous, disinfectant, escharotic, alterative, given in very small doses in syphilitic affections, and in syphilitic and non syphilitic skin diseases. Externally as a **lotion**, 1 grain to the fl oz, or **ointment**, 2 to 8 grains in the oz, in chronic and parasitic skin diseases, and in acne and freckles, a solution of 1 in 1000 is used for syphilitic ulcers, as an ordinary surgical dressing and in obstetric practice 1 in 2000 to 5000 is sufficient, as an **injection**, 1 grain to 8 fl oz, for chronic discharges, such as leucorrhœa and gonorrhœa, and as a **gargle**, 1 grain in 4 fl oz, for ulcerated and syphilitic sore throat, as a **collyrium**, 1 grain in 8 fl oz. For syphilis by **hypodermic injection**, $\frac{1}{30}$ to $\frac{1}{10}$ grain (with Sodium Chloride), in divided portions in the course of the day. As a local application in diphtheria.

An aqueous solution of 1 in 1000 is employed for disinfecting the hands, towels, sponges, etc, in operative surgery, it corrodes surgical instruments. A solution of the same strength is used for washing infected rooms, furniture and other articles, and for soaking infected linen. The solution is often coloured with aniline blue or methyl violet to guard against its being mistaken for water or other harmless fluid.

The disadvantages of Mercuric Chloride as a disinfectant and antiseptic are due (1) to its forming with albumen an inert and insoluble compound, (2) to its corrosive action on metals, and (3) to its being a powerful poison.

To prevent its antiseptic value being destroyed by the formation of an albuminate, five parts of Tartaric or Hydrochloric Acid should be added to each part of Mercuric Chloride.

In France it is legal to supply registered nurses (for obstetric purposes) with a lotion containing 0 025 gramme Mercuric Chloride and 1 gramme Tartaric Acid per litre, also an ointment containing 1 p c in Vaseline —*A J P* '90, 180.

As a disinfectant of enteric or other infectious stools and urine, an equal quantity of a 1 in 500 acidulated solution should be used. They should be thoroughly mixed and left in contact for at least 2 hours before they are finally disposed of.

Recommended for dysentery in India $\frac{1}{17}$ grain every 4 hours.—*L* '89, ii 901

Injection of Corrosive Sublimate solution in hydrocele.—*L* '97, ii 594, in tetanus—*B M J* '97, i 335, in syphilis—*B M J* '96, i 52, and in other forms of tuberculosis—*B M J* '96, ii 71

A handy and trustworthy preparation for use as hypodermic injection in syphilis is—Mercury Perchloride, 1 grain, Glycerin, 1 fl drim, Distilled Water, 1 fl drim. A stable preparation which can be kept for several days or weeks. 10 minims of the solution = $\frac{1}{17}$ grain Mercuric Chloride.—*L* '01, ii 149

Mercuric Chloride and Sodium Chloride (of each 1 g) in 100 c Water 2 cc injected daily for 2 weeks, a day or two after each injection (Hilber)—*M A* '02 42 Great objection to the use of Mercuric Chloride is the pain caused by it, which is often corrected by the addition of 1/2 grain three times a day in the treatment of lues, etc.—*B M J* '01 ii 64

Mercuric salts are the most efficient antiseptics we possess in surgical practice, and it may be argued that the use of Mercuric salts or Mercury when administered internally is to be a counter-indication to a great extent by its antiseptic property. Mercuric Chloride is more valuable as a bactericide in the treatment of scarlet fever.—*B M J* '03, ii 231

With regard to fluid disinfectants, a 1 in 1000 solution of corrosive sublimate with 24 hours' exposure, destroyed all microbes, including the spores of anthrax, and the tubercle bacilli. Anthrax spores were only destroyed with certainty by Mercuric Chloride. Report on the practical experiments on disinfection undertaken for the London County Council.—*L* '02, i 758, *B M J* '02, i 792

$\frac{1}{2}$ grain dissolved in 2½ oz of Water, injected into the pleural cavity, after tapping, in a case of empyema.—*B M J* '03, ii 78

Experiments on the antiseptic action of Mercury. That Mercury Perchloride, Mercury Oxycyanide and Protargol cannot be injected intravenously into rabbits in sufficient strength to produce an antiseptic effect lasting several days.—*L* '03, i 99

Three to five minims of a 5 per cent solution have been successfully given intramuscularly in 40 cases (B M J '04, ii 60). 3-minim doses of the Liqueur of Calomel given to an infant one month old, 5 to 10 minims three times a day for an infant six months old (*L* '04, ii 1405)

In the case of the hands or fingers or surgical operations. A preliminary wash with soap and hot water, followed by rubbing the hands with 50 per cent alcohol 1 in 1000, and then polishing with a dry sterile cloth shown (*B M J* '03, ii 761) to be the most efficient.

One cc of a solution containing Mercury Perchloride 1 gramme Sodium Chloride pure, 10 grammes Distilled Water, 100 grammes, representing a dose of 1 centigramme injected intramuscularly in syphilis (*M P* '06, i 148) the dose may be increased to 2 centigrammes daily for 20 days

Dose— $\frac{1}{32}$ to $\frac{1}{16}$ grain = 0.0022 to 0.004 gramme

Ph. Ger. maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme

Prescribing Notes—*General.* Prescribed in the form of the Liqueur or given in pills well triturated with Milk Sugar and massed with Diluted Glucose.

Compressed Discs are prepared for making an antiseptic solution 1 in 1000, see Not Official

Incompatibles—Alkalis and their Carbonates, Lime Water, Tartar Emetic, Silver Nitrate, Ferric Acetate, Albumen, Soaps. Decoction of Cinchona, Tannin, Alkaline Sulphates. Potassium Iodide converts it into Mercuric Iodide, soluble in excess.

Official Preparations—Liqueur of Hydrargyri Perchloride, and Liqueur of Hydrargyri Flava. Used in the preparation of Hydrargyri Oleas, Hydrargyri Flava, and Hydrargyrum Ammoniatum.

Not Official.—Corrosive Sublimate Discs, Sublimate Wood Wool, Sublimate Gauze, Sublimate Wool, Injectio Hydrargyri Hypodermica, Injectio Sal Alembrothi Hypodermica, Liquor Hydrargyri et Ammonii Chloridi, Lotio Hydrargyri Acetica, Lotio Hydrargyri Perchloridi, Lotio Hydrargyri Perchloridi Acida, Preservative Solution, Poudre de Sublimé Corrosif et d'Acide Tartrique, Sal Alembroth

Antidotes—In poisoning by Corrosive Sublimate, raw eggs should be administered in large quantity, flour with milk may also be given, the stomach should then be washed out or an emetic employed

Foreign Pharmacopœias—Official in Austr and Hung (Hydrargyrum Bichloratum Corrosivum), Belg (Sublimatus Corrosivus), Dan, Norw and Swed (Chloretum Hydrargyricum Corrosivum), Dutch (Chloretum Hydrargyricum), Fr (Bichlorure de Mercure), Ger, Jap, Russ and Swiss (Hydrargyrum Bichloratum), Ital (Bichloruro di Mercurio), Port (Chloreto Mercurico), Mex and Span (Cloruro Mercurico), U S (Hydrargyri Chloridum Corrosivum)

Tests—Mercuric Chloride dissolves in Water, forming a solution which yields with Ammonia Solution a white precipitate, with excess of Hydrogen Sulphide Solution a black precipitate insoluble in Ammonium Hydrosulphide Solution, and in hot diluted Nitric Acid, with Potassium or Sodium Hydroxide Solution a yellow precipitate, with Potassium Iodide Solution a brilliant scarlet precipitate, soluble in an excess of the reagent or in a considerable excess of the Mercuric Chloride Solution. An aqueous solution, when boiled with Copper foil, gives a grey deposit, which assumes a silvery lustre on being rubbed. Potassium or Sodium Hydroxide Solution does not produce a precipitate in a Glycerin Solution, in solutions containing both Glycerin and Potassium or Sodium Hydroxide, Ammonium Hydrosulphide Solution produces no precipitate. With Silver Nitrate it affords a white precipitate, which when filtered and washed is insoluble in Nitric Acid but dissolves readily in Ammonia Solution.

It is officially required to yield 77.28 to 73.8 p.c of metallic Mercury, corresponding to 98.57 to 99.92 p.c of pure Mercuric Chloride, when heated with excess of Lime. The U.S.P. requires that it shall contain not less than 99.5 p.c of pure Mercuric Chloride, but gives no method for its quantitative determination. The P.G. gives neither a requisite percentage nor a method of determination. Mercuric Chloride contains theoretically 73.8 p.c of metallic Mercury.

The more generally occurring impurities are fixed residue, foreign salts, Arsenic, and metals other than Mercury. The specimen should leave, after sublimation, according to the B.P. only a trace of fixed residue, according to the U.S.P. no appreciable residue, and according to the P.G. it must fuse and is then completely volatilised. Foreign salts may be detected in the filtrate, after precipitation of the Mercury with Hydrogen Sulphide as described below under the Hydrogen Sulphide Test. Arsenic, if present, is also precipitated as Sulphide and may be extracted by digesting with Ammonia Solution, the filtered liquid being evaporated to dryness, moistened with 6 drops of Nitric Acid, dried and examined by the Modified Gutzeit's Test. Heavy metals other than Mercury may be detected by treating the Sulphide insoluble in Ammonia Solution with diluted Nitric Acid,

Foreign Pharmacopœias.—Official in Austr, Ger, Ital, Jap and Sweden, 1 and 2 grammes, red. All made with a mixture of equal parts of Mercuric and Sodium Chlorides. Dutch, 1 gramme, blue.

POUDRE DE SUBLIMÉ, CORROSIF ET D'ACIDE TARTRIQUE (*Fr*) —Powdered Mercuric Chloride, 2.5 grammes, Powdered Tartaric Acid, 10 grammes, Solution of Soluble Indigo (5 p c), 10 drops. Mix to a uniform tint, dry, and divide into ten packets. The contents of each packet, when dissolved in a litre of Water, gives a solution 1 in 4000.

LIQUOR HYDRARGYRI ET AMMONII CHLORIDI —Mercuric Chloride, 10, Ammonium Chloride, 10, Tartaric Acid, 10, Distilled Water, *qs* to make 100 —*B P C*

The addition of Tartaric Acid is necessary to prevent precipitation on diluting the solution with ordinary Water —*B P C*

The two Chlorides dissolve in the Water, but immediately after the addition of the Tartaric Acid a precipitate is formed.

The addition of Tartaric Acid to solution of Mercuric Chloride to prevent the formation of insoluble albuminous compounds when applied to animal tissues. —*B M J* '88, 1 148

Dott called attention to the precipitation of the Mercuric Chloride by Tartaric Acid in dilute solution —*P J* '89, 414

LOTIO HYDRARGYRI ACETICA —Mercuric Chloride, 1, Acetic Acid, 75, Glycerin, 75, Alcohol (90 p c), 250, Rose Water, 500 —*Martindale*

To destroy pediculi and detach the ova

Mercuric Chloride, 0.20, Acetic Acid, 8, Glycerin, 8, Alcohol, 27.50, Rose Water, *qs* to produce 100 —*B P C*

LOTIO HYDRARGYRI PERCHLORIDI (1 in 500) —Mercuric Chloride, 1 oz, Water to 500 fl oz. To be diluted with one to ten parts of Water as directed. Usually tinted with fuchsin or methylene blue —*St Thomas's*

This has been incorporated in the *B P C*

LOTIO HYDRARGYRI PERCHLORIDI ACIDA —Mercuric Chloride, 1 oz, Hydrochloric Acid (strong), 25 fl oz, Water to 500 fl oz. This is used only as a disinfectant for excreta —*St Thomas's*

This has been incorporated in the *B P C* under the title *Solutio Hydrargyri Perchloridi Acida*.

SUBLIMATE WOOD WOOL —Pinewood almost in a state of powder, containing $\frac{1}{2}$ p c of Corrosive Sublimate. It is highly absorbent.

SUBLIMATE WOOL (*Ital*) —Absorbent Wool containing 1 of Mercuric Chloride in 400.

SUBLIMATE GAUZE (*Fr*) —Prepared Gauze containing 0.1 to 0.5 p c of Corrosive Sublimate.

INJECTIO HYDRARGYRI HYPODERMICA —Mercuric Chloride, $\frac{1}{2}$ grain, Sodium Chloride, pure, 5 grains, Water to 1 drm.

Dose —4 to 12 minims = $\frac{3}{10}$ to $\frac{1}{10}$ grain in divided portions in the course of one day.

PRESERVATIVE SOLUTION (for Anatomical subjects) —Corrosive Sublimate, 10 grains, Glycerin, 21 fl oz, Methylated Spirit, *qs* to make 80 fl oz. For injection into the femoral artery.

SAL ALÉMBROTH —Mercuric and Ammonium Chloride, $2\text{NH}_4\text{Cl}$, $\text{HgCl}_2 \cdot \text{H}_2\text{O}$, eq 393.32. *Syn* Ammonio-mercuric Chloride, Salt of Wisdom, Sel de la Sagesse ou de la science.

White rhombic prisms or tabular crystals, which readily part with their Water of crystallisation when exposed to dry air.

It contains theoretically 50.5 p c of metallic Mercury.

Solubility —2 in 1 of Water, 1 in $3\frac{1}{2}$ of Alcohol (90 p c), 1 in 1 of Glycerin.

Medicinal Properties —A powerful antiseptic, but it is not so irritating as Corrosive Sublimate. Used in the antiseptic treatment of wounds.

For Hypodermic injection in syphilis, $\frac{1}{2}$ grain dissolved in 10 minims of Water—*B M J '88, 1 905*

Alembroth Gauze, 1 p c, Wool, 2 p c, they are tinted with aniline blue, and as the colour is bleached by purulent discharge, soakage of the dressing is readily noted

Injectio Sal Alembroth Hypodermica—Mercuric Chloride 32 grains Ammonium Chloride 1 lb, Distilled Water 2 fl oz—*Lock*

Dose—10 minims = $\frac{1}{2}$ grain of Sal Alembroth to be used for an injection

Tests—Mercuric Ammonium Chloride when strongly heated its aqueous solution is neutral in reaction. Litmus paper, whilst an aqueous solution of Mercuric Chloride possesses a faintly acid reaction towards that indicator of neutrality. Potassium or Sodium Hydroxide Solution produces a white precipitate in the aqueous solution. It should leave no weighable residue when ignited with free access of air

HYDRARGYRI SUBCHLORIDUM

MERCUROUS OXIDE.

B.P. Syn—CALOMEL, HYDRARGYRI SUBCHLORIDUM, SUBCHLORIDE OF MERCURY

Hg_2Cl_2 , eq. 467.98

FR, PROTOCHLORURE DE MERCURE, GER, QUECKSILBERCHLORUR, ITAL, CHLORURO MERCUROSO, SPAN, CLORURO MERCURIOSO

A heavy, white or whitish, lustrous, tasteless, impalpable powder which should be protected from the light

Solubility.—Insoluble in Water, Alcohol, Ether

Medicinal Properties.—Alterative, indirect cholagogue, purgative, antiseptic, and diuretic

As an alterative it is used in syphilitic affections, chronic skin diseases, and glandular

Useful in chronic jaundice, and in chronic pharyngitis, repeated vomiting, also, in the gastro-intestinal catarrh and diarrhoea of children, for whom the absence of taste renders it convenient.

As a purgative in biliousness, hepatic and cardiac dropsy, apoplexy, gout, cirrhosis, and in congested and torpid liver due to free living

In enteric fever, the stupor, tremor, headache and coma, all of which may be due to intestinal sepsis and ptomaines, are removed, and the entire aspect of the case changed, by 1 to 3 grains of Calomel—The late *Sir W Broadbent*

In hiccup, one grain every hour is often successful. Its uses are numerous as an inflammation, or as a gargle in syphilitic sore throat, as an injection with or without Lime Water, in blenorrhagia. In a wide range of skin affections, but especially syphilitic, it is invaluable as an agent.

For fumigation—A spirit lamp underneath a metal cup containing 20 grains of Calomel is placed under a cane-seated chair on which the patient remains seated for 20 minutes, the chair being covered with a blanket, an apparatus better

It is probable that the cholagogue action of Calomel is due to its having a peculiar stimulant action on the duodenum and ileum, so as to hurry the bile along the intestine and prevent its reabsorption — *Brunton*

Should not be applied to the eye when a patient is taking Potassium Iodide, for it will cause severe inflammation

On the treatment of acute diseases, particularly enteric fever and appendicitis, by a judicious use of Calomel, Water, Heat and Quinine — *B M J '01, ii, 1054*

Although not a specific, is a most useful remedy in typhoid fever — *B M J '99, i 4*

Weekly injections of 0.05 grammes have proved successful (*B M J E '04, ii 73*) in optic neuritis, after injections of the Cyanide and Biniodide have been tried without avail

In enteric fever (*B M J '04, ii 1450*) it has been shown that, of the various drugs which are known to possess antiseptic properties, Calomel is undoubtedly one which has received earliest and widest recognition, 3 to 5 grains are given during the first week of the attack, before there is much diarrhoea

Has been used in large doses in the treatment of dysentery (*I M G '05, ii 280*), 5 to 7 grains every five or six hours, or in smaller doses of 1 grain more frequently. Very frequently prescribed with drugs such as Ipecacuanha and Opium, or in $\frac{1}{2}$ grain doses with 5 grains of Naphthalin 10 or 12 times in the 24 hours. Fractional doses have been given to children

A valuable anthelmintic in ankylostomiasis (*L '05, i 865*)

As an intramuscular injection in syphilis, an emulsion made by the following formula is useful (*M P '06, i 149*) Sublimed Calomel, 1 gramme, pure liquid Vaseline, 10 c c. The average weekly dose is 10 c c

A proteid or colloidal form of Calomel is known under the name of Calomelol. It forms a greyish white odourless and tasteless powder, soluble in Water, but insoluble in Alcohol (90 p c t)

Injections of Calomel the best method of treating syphilis — *L '07, ii 13*

Dose — $\frac{1}{2}$ to 5 grains = 0.032 to 0.32 grammes

Swiss, maximum single dose 0.5 grammes, maximum daily dose, 2.0 grammes

Prescribing Notes — Calomel can be made into pills with Glucose, and if the pills be too small, they can be made larger by the addition of Milk Sugar. It is frequently prescribed with Compound Rhubarb Pill or Compound Pill of Colocynthis and Henbane

Incompatibles — Bromides and Iodides, Nitro-Hydrochloric Acid, Cyanic Acid, Chlorides of the Alkalies, especially when neutral. Sodium Lame, Potassium Hydroxide, or Sodium Hydroxide

Official Preparations — Lotio Hydrargyri Nigra, Pilula Hydrargyri Subchloridi Composita, and Unguentum Hydrargyri Subchloridi

Not Official — Calomel Cream (*Squibb*), Emplastrum Calomelanos, Pastillus Hydrargyri Chlorati cum Talco, Pilula Calomelanos cum Coleo, Pilula Hydrargyri Subchloridi et Jalape, Pilula Hydrargyri Subchloridi et Scammonii, Pilula Zittmann, Pulvis Basilicus, Pulvis Rheum cum Hydrargyro, Pulvis Calomelanos et Acidi Borici, Pulvis Calomelanos et Anthonii, and Pulvis Calomelanos et Zinc Oxidi

For *Ag*n Pharmacopœias — *Offic* in Belg (Calomelas), Dan and Norw (Calomel), Fr (Protochlorure de Mercure par volatilisation), also (Protochlorure de Mercure par Précipitation), Dutch (Chlorretum Hydrargyrosu), also (Chlorretum Hydrargyrosu ope Vaporis Aquæ par steam), Swed (Chlorretum Hydrargyrosu Precipitatum), Austri and Hung (Hydrargyrum Chloratum Mite), both the levigated and that sublimed in steam, Jap, and Swiss (Hydrargyrum Chloratum), also (Hydrargyrum Chloratum Vapore Paratum), Itali (Chloruro Mercurioso), Mex (Chloruro Mercurioso al Vapor), also (Precipitato), Port (Chloreto Mercurioso), also (Marchario Doble), Russ (Hydrargyrum Chloratum Levigatum), also (Hydrargyrum Chloratum Vapore Preparatum), Span (Chloruro Mercurioso) (Sublimado,

por el Vapor, and Precipitado), US (Hydrargyri Chloridum Mite)

The following synonyms are applied to Calomel obtained by precipitation —
Fr., Précipité Blanc Por., Precipitatum Album These terms do not denote Mercury

Tests—Mercurous Chloride volatilises when strongly heated. With Calcium, Potassium or Sodium Hydroxide Solution or with Ammonia Solution it yields a black precipitate, in the case of the three former solutions the precipitate consists of Mercurous Oxide, in the latter case it consists of a Mercurousamido salt. It is converted by Hydrocyanic Acid into a Mercuric salt and a black powder readily yielding metallic Mercury. When heated with alkali Carbonate in a dry test-tube it yields a sublimate of metallic Mercury. If the alkaline residue be dissolved in diluted Nitric Acid it affords with Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid. It is officially required to yield when heated with an excess of Lime, from 84.4 to 84.9 p.c. of metallic Mercury, corresponding to from 99.34 to 99.92 p.c. of pure Mercurous Chloride. The *U.S.P.* states that it should contain not less than 99.5 p.c. of pure Mercurous Chloride but gives no method by which this percentage may be ensured. The *P.G.* gives neither the requisite percentage nor a method of determination. It contains theoretically 84.90 p.c. by weight of metallic Mercury.

The more generally occurring impurities are fixed residue, Mercuric Ammonium Chloride, Mercuric Chloride, foreign salts, e.g., metals of the alkali earths, Arsenic, foreign metals. The *B.P.* requires that when volatilised it shall leave only a trace of fixed residue, the *U.S.P.* no appreciable residue, the *P.G.* that it completely volatilises.

The *B.P.* test for the absence of Mercuric Ammonium Chloride is that the specimen shall not evolve Ammonia when heated with Potassium Hydroxide Solution, the test is supplemented in the *U.S.P.* by the requirement that the filtered Acetic Acid extract shall not be affected by Hydrogen Sulphide or Silver Nitrate Solution. The official method of detecting Mercuric Chloride is by the sample with warm Ether, which on filtration and evaporation should leave no residue. This evaporation must be performed at a low temperature, otherwise the Perchloride (if present) will volatilise in the Ether vapour. The *U.S.P.* requires that the residue after the evaporation of the Ether when dissolved in water shall not yield more than a slight opalescence with Silver Nitrate T.S. and no change in colour with Ammonium Hydrosulphide Solution. The *P.G.* dispenses with an evaporation in testing for the presence of Mercuric Chloride, extracting the sample with Alcohol (60 p.c.), filtering, and requiring that the filtrate shall be unaffected by either Hydrogen Sulphide, or Silver Nitrate Solution. Foreign salts may be detected after the complete removal of the Mercury by Hydrogen Sulphide as described below, the filtrate should leave no weighable residue upon evaporation to dryness and gentle ignition. Arsenic, if present, would be precipitated along with the Mercury as Sulphide and may be extracted by digestion with stronger Ammonia Water and after

treating as described below, examined by the modified Gutzzeit's test. The precipitated Mercuric Sulphide remaining after the Ammonia treatment may be warmed with diluted Nitric Acid and filtered, the filtrate when evaporated to dryness and ignited should leave no weighable residue.

Silver Nitrate—If 1 gramme be shaken with 10 c.c. of dilute Alcohol and filtered, the filtrate should not be affected by T.S. of Silver Nitrate, *P.G.*; 2 grammes of the salt shaken with 20 c.c. of Ether, filtered, the filtrate evaporated and 10 c.c. of distilled Water added, 5 c.c. of the filtrate from this should yield not more than a slight opalescence with Silver Nitrate T.S. A portion of the salt shaken with Acetic A. and filtered, the filtrated liquid should not be affected by T.S. of Silver Nitrate (distinction from and absence of Ammoniated Mercury), *U.S.P.*

Hydrogen Sulphide—If 1 gramme of Mercurous Chloride be shaken with 10 c.c. of Alcohol, as above, the filtrate should not be affected by T.S. of Hydrogen Sulphide, *P.G.*; 1 gramme of the salt shaken with 10 c.c. of Water or Alcohol, and the mixture filtered, the filtrate should not respond to the time limit test for heavy metals, *U.S.P.* The filtrate obtained after shaking a portion of the salt with Acetic Acid and filtering should not be affected by T.S. of Hydrogen Sulphide, *U.S.P.* If to 0.5 gramme of Mercurous Chloride contained in a small beaker, 5 c.c. of Nitric Acid be added, and the mixture evaporated to dryness on a water bath, and if, after dissolving the residue in about 25 c.c. of Distilled Water and 5 c.c. of Hydrochloric Acid, the solution be completely saturated with Hydrogen Sulphide and allowed to stand for several hours in a well corked flask, until the precipitate has subsided, and then filtered, the filtrate should be colourless and leave no weighable residue upon evaporation and gentle ignition, *U.S.P.*

Gutzzeit's Test—The precipitate (obtained in the preceding test) should be washed with about 100 c.c. of Water, then drained and rinsed into a beaker with about 20 c.c. of Water, and then 5 c.c. of Ammonia Solution [sp. gr. 0.897 at 25° C (77° F)] added. After covering this mixture and digesting it for 15 minutes on a water bath, it be rinsed upon a filter and washed with a little Water, the filtrate and washings after evaporating to dryness, moistening with 6 drops of Nitric Acid and again drying, should not respond to the Modified Gutzzeit's test for Arsenic, *U.S.P.*, if the precipitated Sulphide remaining on the filter be treated with diluted Nitric Acid (see an 4), warmed and then filtered, the filtrate should leave no weighable residue upon evaporation and gentle ignition, *U.S.P.*

Ammonium Sulphide.—If 2 grammes of the salt be shaken with 20 c.c. of Ether, filtered, the filtrate evaporated and 10 c.c. of Distilled Water added, 5 c.c. of the filtrate so obtained should not yield any change in colour on the addition of a few drops of T.S. of Ammonium Sulphide, *U.S.P.*

Preparations

LOTIO HYDRARGYRI NIGRA *II.* **BLACK MERCURIAL LOTION**
B.P. Syn—BLACK WASH.

Tinturate 30 grains of Mercurous Chloride with $\frac{1}{2}$ fl. oz. of Glycerin and $1\frac{1}{2}$ fl. oz. of Mucilage of Tragacanth, transfer to a bottle, add 2 fl. oz. of the Solution of Lime, shake well, add sufficient Solution of Lime to produce 10 fl. oz. of the Lotion (about 1 in 146)

Useful application to syphilitic sores, and to relieve itching, as in prurigo senilis and urticaria.

Foreign Pharmacopoeias.—Official *Max* (Agua Fagedenica Negra), 1 in 600. Not in the others.

PILULA HYDRARGYRI SUBCHLORIDI COMPOSITA. COM-
POUND PILL OF MERCUROUS CHLORIDE. *B P Syn* — COMPOUND
CALOMEL PILL, PLUMMER'S PILL

Mercurous Chloride, 1 oz, Sulphurated Antimony, 1 oz, Guaiacum Resin, in powder, 2 oz, Castor Oil, 180 grains, Alcohol (90 p c),
q s about 1 fl drms (1 in 4½)

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

UNGUENTUM HYDRARGYRI SUBCHLORIDI. MERCUROUS
CHLORIDE OINTMENT *B P Syn* — CALOMEL OINTMENT

Mercurous Chloride, 1, Benzoin Lard, 9. (1 in 10)

Useful in the itching of some skin reactions, psoriasis and eczema, also in pruritus ani. A good application to syphilitic sores

Foreign Pharmacopœias—*Offic in Fr* (Pommade de Calomel),
Ital (Pomata di precipitato bianco), 1 in 10, *Port* (Pomada de
Mercurio Doce), 1 in 10, *Mex* (Pomada de Cloruro Mer-
curioso), 1 and 20, *Span* (Pomada de Cloruro Mercurioso
Precipitado), 1 in 10. Not in the others

Not official

CALOMEL CREAM (*Squire*)—Pure hydrosulphurated Calomel, 48 grains,
sterilised anhydrous Lanolin by weight 160 grains, pure sterilised Olive Oil, q s,
to produce 1 oz

10 minims = 1 grain of pure hydrosulphurated Calomel

Dose.—10 minims = 0.6 c c by rectal injection

Calomel, 10 grains, Vaseline to 1—*Lock*

See also Mercurial Cream, p 60

EMPLASTRUM CALOMENOS *Syn* EMPLASTRUM ALBUM—Con-
tains 20 p c of Calomel, spread on or other suitable material

PASTILLI HYDRARGYRI LORATI CUM TALCO (JAP.)—Each
pastil contains 0.5 gramme of Mercurous Chloride

PILULA CALOMELANOS COLOC—Calomel 1, Colocynth
Extract of Colocynth, 3½ grains, Acuanha, ½ fl oz. Compound

Dose.—One or two pills

PILULA HYDRARGYRI SCHLORIDI ET JALAPÆ (House Pill)—
Calomel, 1 grain, Jalap, 3 grains, Syrup of Glucose, q s, in one pill—*St*
Bartholomew's

PILULA HYDRARGYRI SCHLORIDI ET SCAMMONII—Calomel,
1 grain, Scammony, 3 grains, Syrup of Glucose, q s, in one pill—*St*
Bartholomew's

PILULA ZITTMANN—Cal, 2 grains, Compound Extract of Colocynth,
5 grains, Extract of Henbane, 2 grains. Make two pills—*Lock*

Pilulæ Calomelanos Colocynthidis et Hyoscyami *Syn*
Zittmann's Pills—Mercurous Chloride, 1 grain, Compound Extract of Colo-
cynth, 2 grains, Green Extract Hyoscyamus, 1 grain, to make one pill—
B P C

PULVIS BASILICUS—Mercurous Chloride 3, Scammony, 3; Acid
Potassium Tartrate, 3, Jalap-Ginger, 1, Antimonial Powder, 1. Dose for
a child of two years, 4 grains (gramme), of six years or upwards, 8 grains
(0.52 gramme)—*Martindale*

This has been incorporated in the *B.P.C* under the title *Pulvis*
Hydrargyri Subchloridi Compositus.

PULVIS RHEI CUM HYDRARGYRO—Rhubarb Root, in powder, 2 grains, Mercurous Chloride, $\frac{1}{2}$ grain, Ginger, in powder, $\frac{1}{2}$ grain Dose for a child of twelve months—*St Thomas's*

This has been incorporated in the *B P C*

This form is given in *London Ophthalmic* under the title **Pulvis Calomelanos cum Rheo**

PULVIS CALOMELANOS ET ACIDI BORICI—Mercurous Chloride, 1, Boric Acid, in powder, 3 Used as a dusting powder—*St Thomas's*

This has been incorporated in the *B P C*

PULVIS CALOMELANOS ET AMYLI—Mercurous Chloride, 1, Starch Powder, 8 Used as a dusting powder—*St Thomas's*

This has been incorporated in the *B P C*

Mercurous Chloride, 1, Starch, 11—*King's and Lock*

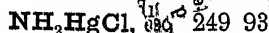
PULVIS CALOMELANOS ET ZINCI OXIDI—Mercurous Chloride, 1, Zinc Oxide, 8 Used as a dusting powder—*St Thomas's and London*

This has been incorporated in the *B P C*

HYDRARGYRUM AMMONIATUM.

AMMONIATE OF MERCURY

B P Syn—AMMONIO CHLORIDE OF MERCURY, MERCURIC AMMONIUM CHLORIDE
WHITE PRECIPITATE



White pulverulent masses or a white odourless powder, possessing a somewhat earthy and subsequently metallic taste. It is converted into a yellow basic salt by prolonged contact with Water

It is known as *infusible white precipitate*

The *fusible* variety is obtained by adding a solution of Mercuric Chloride to a mixture of Ammonium Chloride and Ammonia till the precipitate ceases to redissolve. It has the formula $\text{HgCl}_2 \cdot 2\text{NH}_3$.

Solubility—Soluble in Hydrochloric Acid. Insoluble in Water, Alcohol (90 p c) and Ether

Medicinal Properties—Never given internally. Used in the form of ointment for chronic and parasitic skin diseases, impetigo, herpes, ringworm and scabies. The ointment is used for pediculi, but the powder can be used alone or mixed with Rose Water, and the unpleasantness of greasing the line is avoided.

Official Preparation—Unguentum Hydrargyri Ammoniatum

Not Official—Lowndes' Cream

Antidotes—Stomach-tube or an emetic, preceded by raw eggs and raw flour and water.

Foreign Pharmacopœias—Official in Austr and Hung (Hydrarg. Bichloratum Ammoniatum), Belg (Præcipitatum Album), Dan, Norw and Swed (Chloretum Amido-Hydrargyricum), Dutch (Chloretum Hydrargyrico-ammoniacum), Ger and Jap. (Hydrargyrum Præcipitatum Album), Ital (Cloramiduro di Mercurio), Russ and Swiss (Hydrargyrum Amidato-bichloratum), U.S. (Hydrargyrum Ammoniatum), Ph Lond 1788 (Calx Hydrargyri Præcipitata), and Span, Præcipitatum Album, Ammoniatum.

Tests—Ammoniated Mercury volatilises when heated at a temperature below redness. When boiled with Stannous Chloride Solution it is reduced and turns grey, and produces metallic Mercury. It is completely soluble in warm and in Sodium Thiosulphate Solution. The solution in the latter reagent evolves Ammonia gas, and a precipitate of Red Mercuric Sulphide when boiled, which is changed into the black variety on prolonged boiling. Dissolved in diluted Nitric Acid it affords a scarlet precipitate with Potassium Iodide Solution. When heated with Potassium or Sodium Hydroxide Solution, it assumes a yellow colour and Ammonia gas is evolved. If the mixture be filtered, the filtrate, when acidified with diluted Nitric Acid, yields with Silver Nitrate Solution a white curdy precipitate, insoluble in Nitric Acid, readily soluble in Ammonia Solution. It is officially required to yield, when heated with an excess of Lime, from 78 to 79 p.c. of metallic Mercury, corresponding to 98.0 to 99.3 p.c. of pure Mercuric Ammonium Chloride. The *BP* 1885 required 77.5 p.c., corresponding to 97.4 p.c. of the pure salt. The *USP* requires that it should contain not less than 78 p.c. nor more than 80 p.c. of metallic Mercury, corresponding to not less than 98.0 p.c. nor more than 100.6 p.c. of the pure salt, but gives no method of determination. The *PG* does not refer to either a requisite percentage of metallic Mercury or to the methods of determination. It contains theoretically 79.5 p.c. w/w of metallic Mercury.

The more generally occurring impurities are fixed residue, Mercurous salts, Carbonate, foreign salts, Arsenic and other metals. The *BP* requires that it shall leave on volatilisation only an amount of fixed residue, both *USP* and *PG* require that it shall volatilise without residue. All three Pharmacopœias agree that it should not fuse. The absence of Mercurous salts and Carbonate is ensured by the specimen dissolving completely in Hydrochloric Acid without effervescence. Impurities may be detected by precipitating the Mercury as Sulphide with Hydrogen Sulphide, filtering and evaporating the filtrate to dryness, as described under Hydragryni Subchloridum. The precipitated Mercury Sulphide contains any Arsenic which may have been present, the latter may be extracted by digestion with strong Ammonia Solution, and when treated as described under Hydragryni Subchloridum may be detected by the modified Gutzert's test. Metals other than Mercury, precipitable by Hydrogen Sulphide, may be detected by warming the precipitated Sulphide with diluted Nitric Acid and filtering, the filtrate on evaporation and ignition should leave no weighable residue.

Preparation

UNGUENTUM HYDRARGYRI AMMONIATI. AMMONIATED MERCURY OINTMENT. *BP* *Str.*—WHITE PRECIPITATE OINTMENT. Various of Mercury, 1, Paraffin Ointment, white, 9 (1 in 10)

Foreign Pharmacopœias. Official in Dutch (Ung. Chloreti Hydragrynico-ammonici), 1 in 10; Ger., Jap. and Swiss (Ung. Hydragrynico-ammonici), 1 in 10.

gyri Album), and Russ (Ung Hydrargyri Amidato bichlorati),
1 in 10, U S, 1 in 10 Not in the others

Not Official.

LOWNDES' CREAM —Ammoniated Mercury Ointment, 1, Zinc Ointment, 3, Glycerin, 2, form a cream

HYDRARGYRUM CUM CRETA.

MERCURY WITH CHALK

P B Syn — GREY POWDER

A dull grey powder, free from grittiness, made by thoroughly mixing 1 of Mercury with 2 of Prepared Chalk

Solubility —Insoluble in Water, partially soluble in diluted Hydrochloric Acid, and in diluted Acetic Acid, leaving a greyish residue of finely-divided Mercury

Medicinal Properties —Chiefly given to children as a cathartic, suitable for the prolonged administration of Mercury in syphilis

Half a grain, combined with aromatic chalk powder, can usually be given three times daily in infantile syphilis, and if there is any loosening of the bowels Dover's powder may be given with the grey powder in doses of $\frac{1}{4}$ grain for infants over three months of age, and $\frac{1}{2}$ grain at the age of six months (*L* '04, ii 1405)

Dose —1 to 5 grains = 0.06 to 0.32 gramme

Prescribing Notes —*Best given as a powder by itself, or with Rhubarb, sometimes in cachets, but when required to be made into pills, 'Diluted Glucose' is the best excipient*

Not Official. —Pilula Hydrargyri cum Creta et Opio, Pilula Hydrargyri cum Creta et Ipecacuanha

Foreign Pharmacopœias —Official in Jap and Swed, same as Brit, Mex, Polvo de Mercurio Calcereo, 1 in 23, Part, Mercurio com Carbonato de Cal, 3 in 10, U S, 3 in 10 Not in the others

Tests —Mercury with Chalk, when mixed with Water, acidified with diluted Hydrochloric Acid, and boiled with a bright strip of Copper foil, gives a grey deposit, which assumes a silvery lustre on being rubbed. It partially dissolves in Hydrochloric Acid with effervescence, and the evolved gas, if passed through Lime Water, yields a white precipitate, the resulting solution, when filtered from the deposit of Mercury, should not afford any white or grey precipitate on the addition of Stannous Chloride Solution, indicating the absence of Mercuric salts. The *USP* requires that the solution obtained by digesting 1 decigramme of the powder with 20 c.c of warm diluted Hydrochloric Acid should yield a filtrate, which should be unaffected by Hydrogen Sulphide Solution. A limit of Mercurous oxide is fixed in the *USP* by the requirement that the filtrate obtained when the specimen is digested with warm Acetic Acid and filtered, should not become more than slightly opalescent on the addition of one or two drops of Hydrochloric Acid. The *B.P.* does not specify a necessary method of determination.

Not Official

PILULA HYDRARGYRI CUM CRETA ET OPIO.—*Syn.* Hutchinson's Pills Mercury with Chalk, 1 grain Compound Powder of Ipecacuanha, 1 grain—*St Thomas*

This has been incorporated in the *B P C*

PILULÆ HYDRARGYRI CUM CRETA ET IPECACUANHÆ—Mercury with Chalk, 1 grain, Compound Ipecacuanha Powder, 1 grain—No 1, *University*

HYDRASTIS RHIZOMA.

HYDRASTIS RHIZOMÆ

FR, HYDRASTIS, GER, HYDRASTIS, ITAL, IDRASTE; SPAN, HYDRASTIS

The dried Rhizome and Roots of *Hydrastis Canadensis*, L

Hydrastis contains the alkaloids—Serberine (3 to 4 p c), Hydrastine (2 to 3 p c), and Canadine

The Hydrastis of the *U S P* is required to yield not less than 2.5 p c w/w of Hydrastine, neither that of *B P* nor the *P G* is required to yield a definite percentage of Hydrastine

Medicinal Properties.—Haemostatic, astringent. Useful in chronic catarrhs of the mucous membranes, such as the gastro-intestinal, but especially that of the uterus. Recommended in menorrhagia

The fluid extract is a sovereign remedy as a preventive in spontaneous epistaxis—*MA* '95, 246. It may be used internally or as a 5 p c solution in water as a spray, internally also in aggravated cases of hyperidrosis—*MA* '95, 322. Used locally in chronic pharyngitis—*L* '89, 1 549. 20 to 30 drops of the fluid extract for controlling night sweats—*Pr* 1v 624. In chronic bronchitis—*B M J E* '97, 1 84, '97, 11 60, *Ph* 1x 224

Prescribing Notes.—Equal parts of the Tincture and Water, or 1 of the Liquid Extract to 19 of Water forms a 2.5 p c solution.

Official Preparations.—Extractum Hydrastis Liquidum and Tinctura Hydrastis

Not Official.—Glycerium Hydrastis, Hydrastin, Hydrastina, Hydrastine Hydrochloridum, Hydrastinina, Hydrastinina Hydrochloridum

Foreign Pharmacopœias.—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Ital, Jap, Mex, Norw, Russ, Spain, Swed, Swiss and U S

Descriptive Notes.—Hydrastis Rhizome is of a yellowish-brown colour, and the transverse fracture is hard and short, exhibiting a greenish-yellow, resinous surface in which a row of bright yellow, narrow, distant wood bundles are formed a ring. It has a characteristic odour, and a bitter taste. It is tortuous, simple or branched, $\frac{1}{2}$ to $1\frac{1}{2}$ inches (12 to 38 mm) long and $\frac{1}{8}$ to $\frac{1}{2}$ inch (3 to 12 mm) in thickness, *B P*, sub-cylindrical, oblique, with thin brittle roots, 2 to 5 cm long, and 3 to 6 mm in diameter, with short stem remnants, or stem scars, and slightly annulate. *U S P* It is furnished with numerous brittle slender rootlets, except on the upper surface, which presents short, nearly erect branches, each with terminal cup-shaped scars where a stem has been given off.

Tests—A useful method for the separate determination of Berberine and Hydrastine is given (*YBP* '01, 408, *CD* '01, n 235). A weighed quantity of 10 grammes of the drug in a state of fine powder is exhausted with hot Alcohol, either in a Soxhlet or in a flask fitted with a reflux condenser. The liquid is cooled and the volume adjusted to 100 cc by the addition of Alcohol. A measured quantity of 25 cc of this liquid is placed in a flask of about 8 oz capacity mixed with 1½ cc of Hydrochloric Acid (32 pc), ¼ cc of Sulphuric Acid, and 125 cc of Sulphuric Ether. The mixture is cooled, well shaken, allowed to stand 24 hours in a refrigerator for the crystals of Berberine Hydrochloride to separate, and filtered through a weighed filter paper, the filtrate being reserved. The crystals are washed with a mixture of equal volumes of Alcohol and Ether until the washings cease to give an acid reaction, the washings being added to the main filtrate. The crystals are dried at 105° C (221° F) and weighed. The weight multiplied by 0.9017 and then by 40 gives the percentage of Berberine present in the sample operated on.

The filtrate and washings from the Berberine Hydrochloride crystals are rendered very nearly neutral or only faintly acid, evaporated nearly to dryness on steam-bath, the residue treated with hot Water in small quantities, filtering into a stoppered separator, until the washings from the residue cease to give an alkaloidal reaction with the ordinary test reagents. Sufficient Ammonia Solution is added to the aqueous extract in the separator to render it alkaline and the liberated alkaloid extracted by agitation with Ether. The Ether extraction is repeated until the whole of the Hydrastine is removed, the excess of Ether is removed by evaporation and the alkaloid extracted from its ethereal solution by several agitations with successive portions of a 5 pc Sulphuric Acid Solution. The separated acid liquids are mixed, sufficient Ammonia Solution added to render them alkaline and the Hydrastine extracted by repeated agitation with successive quantities of Ether. The Ether is evaporated, the alkaloidal residue dissolved in an excess of Twentieth-normal Volumetric Sulphuric Acid Solution, and the excess of Volumetric Acid Solution titrated with Hundredth-normal Volumetric Sodium or Potassium Hydroxide Solution. The number of cc of Hundredth-normal Volumetric Sodium Hydroxide Solution used, divided by 5 and subtracted from the number of cc of Twentieth-normal Volumetric Sulphuric Acid Solution employed to dissolve the alkaloidal residue, the product multiplied by 0.019016 and then by 40, yields the percentage of Hydrastine present in the sample.

The above process has been tried in the author's laboratory and found to work well. A sample of the Rhizome gave 3.6 pc of Berberine and 3.20 pc of Hydrastine.

The *U.S.P.* adopts a method for the determination of Hydrastine, which may be briefly outlined as follows—A weighed quantity of 15 grammes of Hydrastis in No. 60 powder is shaken during 10 minutes in a 250-cc. separator flask, with 150 cc of Ether. 5 cc of ammonia solution is added and the flask again shaken at intervals

for half an hour. After the addition of 15 cc of Water to cause the drug to agglomerate, 100 cc of the clear Ether solution is removed to a separator and shaken with 15 cc of Normal Volumetric Sulphuric Acid Solution. The lower acid layer is removed to a second separator, and when the two liquids have separated, the Ether Solution is again shaken with 5 cc of Normal Volumetric Sulphuric Acid Solution and 5 cc of Water, the acid solution being again drawn off, when the liquids have separated. The Ether solution is then shaken with 5 cc of Water, which is in turn removed. The mixed acid and aqueous liquids are mixed with sufficient Ammonia Water to render the liquid alkaline and the liberated alkaloid shaken out with 15 cc of Ether. After separation of the liquids, the lower alkaline portion is drawn off and the Ether solution transferred to a tared flask. The extraction is repeated with two successive portions each of 20 cc and 15 cc of Ether, the alkaline liquid being removed in each case and the ethereal solution transferred to the tared flask. The Ether is removed by evaporation, the alkaloidal residue dried till constant in weight at 100° (212° F), cooled and weighed. This weight multiplied by 10 gives the percentage of Hydrastine.

The ash of Hydrastis varies from 5 to 8 pc and should not exceed 10 pc.

Preparations

EXTRACTUM HYDRASTIS LIQUIDUM. LIQUID EXTRACT OF HYDRASTIS

20 of Hydrastis Rhizome, exhausted by percolation with Alcohol (45 pc), reserving the first 17 and evaporating the remainder to a soft Extract which is dissolved in the first portion, and the whole made up with Alcohol (45 pc) to 20 (1 in 1).

The fluid extracts of the *USP*, *Fr* and *PG* are standardised preparations, that of the *USP* is required to contain 2 pc w/v of Hydrastine, that of the *Fr* others 2 pc w/w.

Dose.—5 to 15 minims = 0.3 to 0.9 cc

Foreign Pharmacopœias.—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and US, all 1 in 1, Fr and Mex have a solid extract. Not in the others.

Shoemaker has used the fluid extract as a stimulant and astringent application in skin diseases.—*L* '85, 11 87.

Tests.—Fluid Extract of Hydrastis has a sp. gr of 1.025 to 1.048, contains about 22 pc w/v of total solids and about 40 pc w/v of Absolute Alcohol.

The method adopted by the *USP* for the determination of the Hydrastine is essentially as follows.—A measured quantity of 10 cc of the Fluid Extract is introduced into a measuring flask of 100 cc capacity, together with 85 cc of Water containing in solution 2 grammes of Potassium Iodide. After the addition of sufficient Water to bring the volume of the liquids to 100 cc, they are shaken for several minutes. A filtered measured quantity of 50 cc is rendered alkaline with Ammonia Water, and the liberated alkaloid shaken out with 50 cc of Ether, repeating the extraction with a further quantity

of 20 c c of Ether. The lower alkaline liquid is in each case removed after complete separation, the Ether solution transferred to a tared flask, the mixed ethereal solutions evaporated at a gentle heat, the residue dried on the water-bath till constant in weight, cooled and weighed. This weight multiplied by 20 gives the percentage w/v of Hydrastine. The *USP* process works satisfactorily, and the alkaloid is obtained in a condition of purity. A sample of the *BP* liquid extract prepared and examined in the author's laboratory had a sp gr of 1.048, contained 22.2 p c w/v of total solids and 35.4 p c w/v of Absolute Alcohol. When assayed by the *USP* process, it yielded 2.12 p c w/v of Hydrastine.

The essential details of the *P* method for the determination of the Hydrastine content are as follows.—A weighed quantity of 15 grammes of the Fluid Extract is evaporated on a water-bath to about 5 grammes, the residue transferred to a flask, 10 c c of Petroleum Ether, 50 grammes of Ether, and 5 grammes of Ammonia Solution added, and the mixture allowed to stand, with frequent intervals of shaking, for one hour. A weighed quantity of 50 grammes of the clear Ether solution is filtered into a separator, 10 c c of a mixture of 1 part of Hydrochloric Acid and 4 parts of Water added, the whole is shaken for a few minutes. After complete separation the acid layer is drawn off into a beaker, the ethereal solution is washed twice in succession with 5 c c of Water, to which a few drops of Hydrochloric Acid have been added, these washings being added to the main quantity in the beaker. The mixed acid liquids are rendered alkaline with Ammonia Solution, 50 grammes of Ether are added and the mixture allowed to stand for one hour, shaking frequently at intervals. A weighed quantity of 40 grammes is filtered through a dry filter paper into a dry tared flask, the Ether distilled off, and the residue dried at 100° C. (212° F), cooled and weighed. It should amount to at least 0.2 gramme, corresponding to 2.0 p c w/v of Hydrastine.

TINCTURA HYDRASTIS. TINCTURE OF HYDRASTIS

2 of Hydrastis Rhizome in No. 60 powder, percolated with Alcohol (60 p c) to yield 20

Dose — $\frac{1}{2}$ to 1 fl dr = 18 to 36

Foreign Pharmacopœias—Official in Ger., and Ital., 1 and 5, Span., 1 in 10, *US*, 1 in 5. Not in the others.

Tests.—Tincture of Hydrastis has a sp gr of 0.920 to 0.930, contains about 2.5 p c w/v of total solids and about 58.0 p c w/v of Absolute Alcohol. The *BP* Tincture is not a standardised preparation. The *USP* 1 in 10 Tincture is required to contain 0.40 p c w/v of Hydrastine. The *PG* does not include a Tincture. The method of determination of the Hydrastine adopted by the *USP* is essentially as follows.—A measured quantity of 100 c c is concentrated by evaporation to about one-tenth its volume, sufficient Alcohol (94.9 p c) being added to dissolve any insoluble matter which may possibly have been present. The same method of determination is adopted for the assay of the Tincture as for the assay of Fluidextractum.

page of Hydrastine it must be taken into account that the weight of residue represents that from 50 c.c. of the Tincture, and must therefore be multiplied by 2 and not by 20 as in the case of the Fluid Extract prepared in the author's laboratory. A specimen of *B P* Tincture prepared by the process described above, yielded, when assayed by the *U.S.P.* method, 0.288 g w/v of Hydrastine.

Not Official. — Ten 100 of Hydrastis with 85 of

GLYCERITUM HYDRASTIS — Macerate enough Alcohol (95 p.c.) to saturate the powder and leave a stratum above it, macerating for 48 hours, continue percolation with more Alcohol until the Hydrastis is practically exhausted. Remove nearly all the Alcohol by distillation or evaporation in a cold place, and add liquid into 50 of ice-cold Water, and set it aside to make the filtrate. Dose, 30 minims. — *U.S.P.* of Glycerin and mix thoroughly. Average *B.P.C.* under the title **Glycerinum Hydrastis**, with the *syn* Glycerinum Hydrastis.

This has been incorporated in the *U.S.P.* Hydrastis, with the *syn* Glycerinum Hydrastis.

HYDRASTIN — An eclectic remedy, formerly prepared from Hydrastis. It is an extract in fine powder with Alcohol (90 p.c.). Care must be taken not to confound it with Rhizoma Hydrastis.

Dose — 2 to 6 grains = 0.13 to 0.39 grammes.

Considerable variation has been shown (*C.D.* '01, p. 235) to exist among commercial specimens of Hydrastis. In 10 samples were examined, four American specimens, one manufactured by a reliable firm, by the *B.P.C.* (1901) process, and one from an unknown source. The percentage of Hydrastine varied from 17.2 p.c. to 26.9 p.c. in the remaining specimens, and trace in four of the preparations to 8 p.c. The preparations to 9.67, 16.53, 23.5, and the Berberine from a trace in two of the preparation made by the *B.P.C.* (1901) 26.9 p.c. in the remaining four. The following suggestions are made for the standardisation of Hydrastine. The extraction and the product be that a stronger Alcohol be used for Hydrastine. The two former suggestions have been adopted in the *B.P.C.* (1907), but no standard is adopted, though a note at the end of the monograph mentions 1.20 p.c. of total alkaloids, of which two-fifths should be Hydrastine.

HYDRASTINA Hydrastine $C_{21}H_{21}NO_6$, eq. 380.33 — An alkaloid crystallising in white glistening four-sided prisms.

It is obtained from the Rhizome of Hydrastis Canadensis. A close relationship exists between Hydrastine and Narcotine. Schmidt considers that Narcotine contains three Methoxyl groups and Hydrastine only two. Both bases yield on oxidation one such group, it therefore follows that Covarine contains only one and that Cotarnine has the constitution of a Methyl-hydrastine.

Solubility — 1 in 120 of Alcohol (90 p.c.), 1 in 83 of Ether, 1 in 2 of Chloroform. The last two solvents do not dissolve Berberine, insoluble in Water and Petroleum Ether.

Dose — $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.016 to 0.032 grammes. Phosphate forms a soluble compound which can be made to contain 71 p.c. of Hydrastine. — *A.J.P.* '97, 604, *P.J.* '98, 194. — Official in Fr. and U.S.

Foreign Pharmacopœias — 182° C. (209° F.). It possesses an alkaline reaction. Tests — Hydrastine melts at 182° C. (209° F.). It possesses an alkaline reaction towards moistened red

Official Names in Roman; all others in Italics

EUD

	Dose	Page		Dose	Page
<i>Erythimum Bonjean Depuratum pro Iniectione</i>	1 to 4 gr	490	<i>Essential Oil of Camphor</i>		312
" <i>Denzel Fluidum</i>	3 to 10 gr	490	<i>Essenza de Amica</i>		164
" <i>Kohlmann Fluidum</i>	60 to 75 gr	490	" <i>de Cayeput</i>		278
" <i>Purum Dialysatum Wer nich Fluidum</i>	10 to 60 gr	490	" <i>de Cannella</i>		397
" <i>Purum Dialysatum Wer nich Spissum</i>	10 to 30 gr	490	" <i>Cedro</i>		723
" <i>Purum Dialysatum Wer nich Succum</i>	22 gr	490	" <i>Garofani</i>		334
" <i>Purum Succum Wiggers</i>	1 to 1 gr	491	" <i>Ginepro</i>		706
<i>Eigotomae</i>		489	" <i>Lavanda</i>		719
" <i>Hydrochloride</i>		490	" <i>Menta</i>		764
" <i>Osculate</i>		490	" <i>Rose</i>		1023
" <i>Phosphata</i>		490	" <i>Rosmarino</i>		1026
<i>Eriogonon Canadensis Oleum</i>	5 to 10 min	491	" <i>Ruta</i>		1029
<i>Erythrol</i>		25	" <i>Sandalo</i>		1041
" <i>Tetranitrate</i>	1 to 1 gr	491	" <i>Trementina</i>		1198
<i>Erythrophloeum</i>		492	<i>Essig</i>		9
<i>Erythrophloeum Hydrochloridum</i>		492	<i>Essigather</i>		107
<i>Erythrophloeum</i>		492	<i>Essigrose</i>		1021
<i>Essencia de Anis</i>		1062	<i>Essigsauere</i>		9
" <i>Canela</i>		184	<i>Estafisagra</i>		1163
" <i>Cayeput</i>		397	<i>Ester value of Gums and Gum resins</i>		1318
" <i>Clavo</i>		278	<i>Estoragae Liquido</i>		1170
" <i>Espelegio</i>		134	<i>Estramomo</i>		1164
" <i>Lemon</i>		719	<i>Estratto de Felce Maschio Ferezo</i>		540
" <i>Menta</i>		764	<i>Estrofanto</i>		1161
" <i>Piperita</i>		764	<i>Eter</i>		103
" <i>Romero</i>		1026	" <i>Acetico</i>		107
" <i>Sandalo</i>		1041	" <i>Ambintroso</i>		156
" <i>Trementina</i>		1198	" <i>Bombydraco</i>		109
<i>Eserina Sulfato</i>		889	" <i>Nitroso Alcoholizado</i>		1140
<i>Eserina</i>		891	" <i>Sulfurico Alcoholizado</i>		106
" <i>Sahcylate</i>		891	" <i>Yodhidrico</i>		112
" <i>Sulphate</i>		889	<i>Etere</i>		103
<i>Esperma</i>		356	" <i>Acetico</i>		107
" <i>de Ballena</i>		355	" <i>Nitroso Officinale</i>		1140
<i>Espermaceu</i>		356	<i>Ether</i>		103
<i>Espirito d Alecrim</i>		1029	" <i>See also Ether</i>		107
<i>Espirito de Nitro Dulce</i>		1140	" <i>Acetico</i>		107
<i>Espirito Ammoniacal Aromatico</i>		142	" <i>Alcohol Solution of Theobroma</i>		1207
<i>Essences</i>		824	" <i>Excerpt for Tablets</i>		106
<i>Essence de Anis</i>		164	" <i>Alcoolice</i>		111
" <i>de Bugade</i>		203	" <i>Hydrodic</i>		109
" <i>Cannelle de Ceylan</i>		897	" <i>Hydrobromu</i>		106
" <i>Citron</i>		792	" <i>Methylated</i>		106
" <i>d Eucalyptus</i>		495	" <i>Methylic</i>		104
" <i>de Genevre</i>		706	" <i>Purified</i>		1054
" <i>Guayle</i>		334	" <i>Soap</i>		103
" <i>Lavande</i>		719	" <i>Sulphure</i>		107
" <i>Menthe Poiree</i>		704	<i>Ethyl Acetate</i>		100
" <i>Portugal</i>		203	" <i>Bromade</i>		1225
" <i>Rosmarin</i>		1026	" <i>Carbamate Ethyl Urethane</i>		110
" <i>Rose</i>		1023	" <i>Chloride</i>		1225
" <i>Rue</i>		1029	" <i>Ester</i>		104
" <i>Santal</i>		1041	" <i>Hydroxide</i>		111
" <i>Terebinthine Officinale</i>		1198	" <i>Iodide</i>		112
" <i>of Ginger</i>		1257	" <i>Nitritus Liquor</i>		1142
<i>Essencia</i>		873	" <i>Oxide</i>		103
" <i>de Alecrim</i>		824	<i>Ethy ene ethenyl diamine</i>		905
" <i>Alfasema</i>		1027	" <i>Perodula</i>		665
" <i>Horlela</i>		720	<i>Eucalyptine (A)</i>		413
" <i>Pimenta</i>		763	" <i>(B)</i>		413
" <i>Linnao</i>		765	" <i>Hydrochloride (A)</i>		413
" <i>Noz Moschada</i>		723	" <i>(B)</i>		413
" <i>Rosa</i>		799	" <i>Lactate</i>		415
" <i>Ambro</i>		1023	<i>Eucalyptene Bichloride</i>		498
<i>Essencia Anisi</i>		707	<i>Eucalyptol</i>		493
" <i>Camphore</i>		166	<i>Eucalypti Gummi</i>		492
" <i>Citri</i>		812	<i>Eucalyptum</i>		494
" <i>Mentha Piperita</i>		723	" <i>Oleum</i>		498
" <i>Myristica</i>		790	<i>(Eucalyptol Crystallizable)</i>		499
<i>Essentia</i>		521	" <i>Phosphate</i>		498
			<i>Eucalyptus Gauze</i>		498
			" <i>Kumut</i>		711
			" <i>Linal</i>		498
			" <i>Wol</i>		498
			<i>Eucalyptus</i>		983
			<i>Fuchlon</i>		970
			<i>Euderm</i>		1190

laevogyrate, whilst the solution in diluted Sulphuric Acid is dextrogyrate. It dissolves in Sulphuric Acid without change of colour in the cold, but on heating a purple violet colour is produced. Sulphuric Acid containing a trace of Molybdic Acid (Froehde's reagent) produces at first a green coloration, subsequently changing to brown. Sulphuric Acid containing a trace of Selenous Acid yields a yellowish-red colour, changing to a brown. Sulphuric Acid containing a trace of Nitric Acid gives a yellow to orange red colour. Pure Nitric Acid yields an orange-coloured solution, depositing an insoluble substance on the addition of Water, the liquid exhibiting an intense blue fluorescence. When dissolved in Sulphuric Acid and brought into contact with a clear crystal of Potassium Bichromate a red colour changing to brownish is produced. The solution obtained by dissolving a crystal of the alkaloid in diluted Sulphuric Acid instantly decolorises Potassium Permanganate Solution, and an intense blue fluorescence is developed. An excess of Potassium Permanganate Solution must be avoided. It may be distinguished from Hydrastinine by the Potassium Permanganate Test described above. It may be distinguished from Berberine by giving no red coloration with Chlorine Water. It may be distinguished from Strychnine and Gelsemine by the instantaneous appearance of a bright red colour when a small portion of the precipitated Bichromate is touched with a drop of Sulphuric Acid. It should leave no weighable residue when ignited with free access of air.

HYDRASTINÆ HYDROCHLORIDUM Hydrastine Hydrochloride $C_{21}H_{21}NO_3$, HCl, eq 416.52—Pale yellow semi-crystalline powder.

It is very hygroscopic and should, therefore, be kept in well-closed glass bottles, preferably of an amber tint, in a cool atmosphere, and exposed as little as possible to contact with the air.

It contains theoretically 91.3 p.c. of Hydrastine.

Solubility—About 1 in 1 of Water and about 1 in 1 of Alcohol (90 p.c.).

Dose— $\frac{1}{2}$ to 1 grain = 0.032 to 0.065 gramme.

Has been used as an ecboic to induce premature labour, maximum daily dose, $\frac{7}{8}$ grains internally, 5 grains by hypodermic injection—*L* '86, 1990, its physiological action—*B M J* '98, 11 1052.

Tests—Hydrastine Hydrochloride melts at about $116^{\circ}C$ ($240.8^{\circ}F$). Its aqueous solution is neutral in reaction towards Litmus. The aqueous solution yields with Potassio-mercuric Iodide (Mayer's) Solution an amorphous yellowish-white precipitate, with Iodo-potassium Iodide (Wagner's) Solution, a deep brown flocculent precipitate, with Picric Acid Solution, a yellow amorphous precipitate, with Potassium Bichromate Solution, a yellow precipitate, soluble in excess of the reagent, with Potassium Ferrocyanide, a yellow precipitate, soluble in excess of the reagent. Hydrastine may be obtained by shaking out the aqueous solution rendered alkaline with Ammonia Solution, with Ether. The residue left on the evaporation of the Ether should respond to the tests given under Hydrastina. It should leave no residue when ignited with free access of air.

HYDRASTININA Hydrastinine $C_{21}H_{21}NO_3$, eq 205.59—This formula is that given in the *Fr Codex* and in the best known text books. The formula $C_{21}H_{21}NO_3$ appears in the *B.P.C*.

An oxidation product of the natural alkaloid Hydrastine. Colourless or light yellow crystals. Not readily soluble in Water, but soluble in Alcohol, Ether and Chloroform.

Foreign Pharmacopœias.—Official in *Fr* and *Mex*.

Tests—Hydrastinine has a m.p. of 116° to $117^{\circ}C$ (240.8° to $242.6^{\circ}F$). With the majority of the mineral acids it forms salts soluble in Water. Its solution in diluted Hydrochloric Acid is optically inactive, it possesses a faint fluorescence and a very bitter taste. It may be distinguished from most other alkaloids by its powerful reducing action upon Potassio-mercuric Iodide (Nessler's) Solution, an immediate black precipitate being thrown down. Among the alkaloids, Morphine and Apomorphine appear to be the only ones acting in a similar manner, and among the glucosides, Picrotoxin. It should leave no weighable residue when heated with free access of air.

HYDRASTINÆ HYDROCHLORIDUM Hydrastine Hydrochloride

C.H., NO, HCl, eq 223 90 —Light yellow needle-shaped crystal, or a pale yellow crystalline powder Soluble in its own weight of Water, 1 : 30. Alcohol (90 p c)
It should be kept in well-closed bottles

It contains theoretical 88.8 p c of anhydrous Hydrastinine

Medicinal Properties —Useful in endometritis, and uterine fibroid, in which excessive bleeding is a prominent symptom —*L* '90, 1 712, *T G* '90, 86, '92, 539, 699, *Pr* xlv 378 Valuable in menorrhagia —*L* '92, 11 1350, *L* 94, 1 1521

Checks uterine hemorrhage, ameliorates night sweats in phthisis During labour it undoubtedly strengthens feeble contractions and revives an inert uterus. —*B M J E* '93, 1 63

Dose — $\frac{1}{2}$ to 1½ grains = 0.032 to 0.100 gramme, used hypodermically in a 10 p c aqueous solution

Ph G. maximum single dose, 0.03 gramme, maximum daily dose, 0.1 gramme

Foreign Pharmacopœias —Official in Belg, Fl, Ger, Swiss and U S Not in the others

Tests —Hydrastinine Hydrochloride melts at 212° C (413.6° F) The aqueous solution is neutral in reaction towards Litmus paper, it exhibits a strong blue fluorescence, especially when highly diluted An aqueous solution affords with Bromine Water a yellow precipitate soluble to a clear solution in Ammonia Solution, with Potassium Bichromate a yellow precipitate soluble on warming the solution but again insoluble in crystals when the solution cools Ammonia Solution to which 1 drop of turbidity 4 or 5 drops of Sodium Hydroxide Solution (15 p c) addition of 0.1 gramme of the salt dissolved in 3 cc of Water produce after each addition a white turbidity, on shaking on continued shaking or on stirring with a glass rod, the turbidity disappears, the solution becomes clear, the crystalline Hydrastinine separates on the supernatant liquid shall be perfectly clear and not more than a pale yellow colour

Sulphuric Acid produces a deep yellow colour, Sulphuric Acid with a trace of Nitric Acid a reddish-brown colour, and Nitric Acid produces a deep yellow colour with the salt It should leave no weighable residue when dried with free access of air

Not Official

HYDROGENII PEROXIDUM.

H₂O₂, eq 33 76

In its purest condition this is a colourless liquid, sp gr 1.452, evolving when heated 475 times its volume of Oxygen gas It is obtained by decomposing Barium Peroxide with Sulphuric Acid, and concentrating the solution in vacuo over Sulphuric Acid Commercially it is sold containing or available Oxygen

Perhydrol is stated to be a chemically pure Hydrogen Peroxide which, although it reddens Litmus strongly, is free from Acids

Official Preparation

LIQUOR HYDROGENII PEROXIDI. SOLUTION OF HYDROGEN PEROXIDE.

A colourless, almost odourless liquid possessing a slightly acidulous taste, and producing a peculiar sensation on and about the mouth It is an aqueous solution of Hydrogen Peroxide containing 9 to 11 volumes of available Oxygen, equivalent to about 3.0 p c by weight.

It may be prepared by the interaction at a temperature below 10° C (50° F.) of Barium Peroxide Water and a dilute mineral acid

It appears as Aqua Hydrogenii Peroxidi in the U.S.P.

Medicinal Properties—It parts with its Oxygen freely, and is a most powerful oxidising agent and disinfectant. It is a non-poisonous antiseptic. It does not precipitate albumen, and does not interfere with the action of Pepsin, Pancreatin, or Malt Extract. Used locally as a surgical dressing and for purulent discharges, and as a spray or swab in diphtheria. A spray of 10 volume strength is a good application to the throat in scarlet fever, and a 5 volume solution as a deodorising gargle. It is used for bleaching hair and delicate fabrics. It has been recommended internally in enteric fever, chronic bronchitis, and diabetes. It is not well adapted for hypodermic injection, because of the gas it evolves, although in cases of cyanide poisoning it is worth the risk of emboli.

Rapid healing of chancres by spray. — *M A* '95, 168

As a spray in the treatment of *Infus vulgaris* and tubercular abscess — *B M J* '02, 1 448

As a wash in the treatment of suppurative lesions of the skin — *T G* '01, 689.

Injectons of Oxygenated Water diluted with five times its volume of warm sterilised Water, and preceded by an evacuant injection, in the treatment of infantile dysentery — *L* '02, 1 392

A bandage soaked with solution of Hydrogen Peroxide and allowed to dry on the wrist gave rise to spontaneous combustion. This was, no doubt, due to the solution containing Sulphuric Acid. The acid now used for its preservation is Phosphoric.

Dose — $\frac{1}{2}$ to 2 fl drms = 1 8 cc to 7 1 cc

Should be well diluted

Prescribing Notes—*Solution of Hydrogen Peroxide does not keep well, but is liable to lose Oxygen even to the extent of half its strength in a year. Phosphoric Acid is the best preservative and is now generally added for that purpose. When gently warmed it gives off Oxygen very readily. Alcohol and Ether have been used to preserve it, and a solution in Ether is sold under the name Ozonic Ether, the usual strength of which is about equal to 4 volumes of Oxygen.*

Foreign Pharmacopœias—Official in Austr and Swiss (Hydrogenium hyperoxydatum Solutum), Belg and Mex (Agua Oxigenada), sp gr 1 452, Span (Agua Oxigenada), US (Aqua Hydrogenii Dioxidii). All contain 10 volumes of available Oxygen. Fr (Soluté Officiel d'Eau Oxygénée), Ital (Acqua Ossigenata), 12 volumes.

Tests—Hydrogen Peroxide Solution has a sp gr of 1 014. At the ordinary temperature or more quickly when heated it evolves Oxygen. When mixed with an acidified Solution of Potassium Iodide, Iodine is instantly liberated. When mixed with Potassium Permanganate Solution acidified with Diluted Sulphuric Acid a brisk evolution of Oxygen ensues, the Permanganate at the same time being decolorised. A blue coloration appears at the junction of the two fluids, when a few drops of the Peroxide Solution are agitated with 10 cc of Water containing 10 drops of diluted Sulphuric Acid Solution and a drop of Potassium Chromate Solution and a few drops of Ether, the Ether also, after shaking, assuming a blue colour. It is officially required to afford at normal temperature and pressure not less than 18 nor more than 22 volumes of Oxygen, corresponding to about 3 0 p c w/w of absolute Hydrogen Peroxide. The *B P* method of determination is a gasometric one, carried out with a mixture of 2 volumes of 5 p c Potassium Permanganate Solution,

one volume of Sulphuric Acid and 7 of Water. When 1 volume of the Peroxide Solution is shaken with 10 to 12 volumes of such a mixture the quantity of Oxygen evolved above should be liberated, the gas is measured at the normal temperature and pressure. In the reaction which ensues, double the volume of Oxygen is evolved, half being derived from the Peroxide and half from the Sulphuric Acid Solution. The Peroxide method, gasometrically applied, has been subject to some unfavorable criticism and is generally considered as in all cases unreliable.

A suggestion has been made ^{CD '01, ii 222, PJ '01, ii 131, JCS Abs, '01, ii 686} to use a saturated Magnesium Sulphate Solution in the place of the Barium Solution, officially recommended, the latter having been proved to evolve Chlorine in the presence of Sulphuric Acid and Potassium Permanganate Solution, and to thus account for the discordant results obtained by the official process. Confirmation of the accuracy of the Kingzett Titration process given below is also yielded.

The *USP* employs the Potassium Permanganate method volumetrically as mentioned below. Each cc of Tenth-normal Volumetric Potassium Permanganate Solution corresponding to 0.1 p.c w/w of absolute Hydrogen Peroxide or 0.329 volume of Oxygen. The *USP* requirements are that to be of full strength it shall contain 9.87 volumes of Oxygen, corresponding to 3.0 p.c w/w of absolute Hydrogen Peroxide. The method simple, rapid and accurate, which has been used extensively in the author's laboratory, is that of Kingzett. Its accuracy is, moreover, not lessened by the presence of the usual preservative agent. A measured quantity of 10 cc of the Peroxide is mixed with 10 cc of a diluted Sulphuric Acid (1.3 sp gr) and made up to 100 cc with Water. A measured quantity of 10 cc of this solution is then run into 10 cc of a 10 p.c Potassium Iodide Solution, the mixture allowed to stand for 5 minutes and titrated with Volumetric Sodium Thiosulphate Solution. 1 c.c. of the Volumetric Thiosulphate Solution is equivalent to 1.118 cc of Oxygen, but the figure must be divided by 2 to ascertain the number of volumes of available Oxygen.

The more generally occurring impurities are Barium, solid residue, excess of free acid, Arsenic, heavy metals, Hydrofluoric Acid. The *BP* recognises Barium, Arsenic, solid residue as likely impurities, but includes no tests for the detection of these substances. It is officially required to yield no characteristic reaction with the tests for Barium, and to yield when evaporated the residue on a water-bath not more than 0.5 p.c of solid residue. The *USP* requires that no turbidity or precipitate should be produced on the addition of a few drops of diluted Sulphuric Acid to 15 p.c of the Peroxide Solution, that it shall leave not more than 0.5 p.c w/v of total solids as determined by evaporating 20 cc of the solution upon a water-bath and drying the residue at 120° C. The *USP* fixes the limit of free acids at 0.048 p.c w/v, and states that solutions which have been held in stock some time show a marked increase in the percentage w/v of acid. Arsenic may be detected by evaporating

the Peroxide Solution with an equal volume of Ammonia Solution and applying the modified Gutzeit's test, heavy metals and Hydrofluoric Acid by the tests described below. The inclusion of a test for the latter acid in the *BP* has been recommended.

The solution is not official in the *PG*.

An investigation (*CD* '99, 11 213, 240, *PJ* '99, 11 100, 115) showed that the most suitable bottles for the preservation of Hydrogen Peroxide Solution were champagne bottles and soda-water bottles. The effect of such shaking and vibration as specimens would be subjected to in daily transit is also reported upon. Any angular inequality of a bottle hastens decomposition, as also do the small rough patches of Iron Oxide frequently found on stone bottles. If kept in corked bottles, care should be taken that the bottles are not laid on their side, as contact with the cork hastens decomposition.

Gutzeit's Test—The residue obtained on evaporating to dryness on a water bath a mixture of 1 c.c. of the solution and 1 c.c. of Ammonia Water should not respond to the modified Gutzeit's test for Arsenic, *USP*.

Time-limit Test—The residue obtained by evaporating 1 c.c. to dryness on a water bath, when dissolved in a mixture of 9 c.c. of Water and 1 c.c. of diluted Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*.

Sodium Hydroxide and Sulphuric Acid—If 50 c.c. of the solution be rendered alkaline by Sodium Hydroxide T.S. and evaporated to dryness on a water bath, and the residue transferred to a watch glass, moistened with Sulphuric Acid, and allowed to stand for a few hours in a moderately warm place, the surface of the glass after being washed should show no sign of corrosion, indicating the absence of Hydrofluoric Acid, *USP*.

Volumetric Determination of Free Acid—If to 25 c.c. of the solution 5 c.c. of Tenth normal Volumetric Potassium Hydroxide Solution be added and the mixture be evaporated to about 10 c.c. and 3 drops of Phenolphthalein T.S. be added, not less than 2.5 c.c. of Tenth-normal Volumetric Solution of Sulphuric Acid should be required to discharge the red colour of the solution after continued boiling, *USP*.

Volumetric Determination—A measured quantity of 10 c.c. of the solution is diluted with sufficient Distilled Water to measure 100 c.c. A measured quantity of 16.9 c.c. of this liquid is transferred to a beaker, mixed with 5 c.c. of diluted Sulphuric Acid, and Tenth normal Volumetric Potassium Permanganate Solution added from a burette with constant stirring until a faint pink tint is just retained. Each c.c. of the Tenth normal Volumetric Solution represents 0.1 p.c. absolute Hydrogen Dioxide or 0.329 volume of Oxygen. If the solution be of full strength, 30 c.c. of Tenth-normal Volumetric Potassium Permanganate Solution will be required, *USP*.

Not Official.

GUTTÆ HYDROGENII PEROXIDI.—1 Hydrogen Peroxide 10 volumes —
Throat Two or three drops to be poured into the ear, for fetid discharges.

Not Official.

HYGROPHILA.

The dried Herb and Root of *Hygrophila spumosa*, T. And., is official in the *Ind.* and *Col. Add.* for India and the Eastern Colonies.

HYOSCYAMI FOLIA.

HYOSCYAMUS LEAVES.

B P Syn — HENBANE LEAVES

FR, JUSQUIAMI NOIRE, GER, BILSENRAUTBLÄTTER, ITAL, GUSQUIAMO,
SPAN, BELENO

The fresh Leaves, Flowers and Branches of *Hyoscyamus niger*, L.; also the Leaves and the Flowering Tops, separated from the branches and carefully dried. Collected from the flowering biennial plants.

The dried leaves and flowering tops collected from plants of the second year's growth are alone official in the *USP*. The official leaves are not required to yield definite percentage of mydriatic alkaloids, those of the *USP* are required to yield not less than 0.08 p.c. The herb is official in the *P.G.*, but no definite percentage of mydriatic alkaloids is required.

Medicinal Properties.—Sedative, antispasmodic. Similar in action to Belladonna and Scamellaria but milder. Used in insomnia when Opium, from its soporific and other objectionable properties, is not advisable. It is employed to diminish pain and allay irritability of the bladder and to prevent the griping of purgative medicines while it increases the peristaltic action, in visceral neuralgias and in asthma and all spasmodic affections, to allay the irritation of teething and prevent convulsions. Children bear Hyoscyamus well, the aged not so. In large doses it dilates the pupil. Hyoscine is much employed in maniacal delirium.

As a hypnotic it is given in the usual way (*B.M.J.* 1905, ii 1005) to its alkaloid Hydrobromide, which is practically devoid of sleep-bringing properties.

Ph. Ger. maximum single dose, 0.4 gramme, maximum daily dose, 1.2 grammes.

Incompatibles.—Vegetable acids, Silver Nitrate, Iodine, Liquor Potassæ or Sodæ.

Official Preparations.—Extractum Hyoscyami Viride, Succus Hyoscyami, and Tinctura Hyoscyami, used in the preparation of Hyoscine Hydrobromidum, and Hyoscyamina Sulphas. The extract is contained in Pilula Colocynthis et Hyoscyami.

Not Official.—Hyoscyami Radix, Chloroformum Hyoscyami, Huile de Jusquame Composée, Linimentum Hyoscyami, Oleum Hyoscyami, Oleum Hyoscyami Compositum, Oleum Hyoscyami Infusum, Constant Tincture of Hyoscyamus (*Squire*), Tinctura Hyoscyami Radicis, Hyoscyamina, and Hyoscyamine (Scopolamine).

Antidotes.—The same as for Belladonna.

Foreign Pharmacopœia.—Official in Austr., Dutch, Hung., Ital. (Giusquiamo), Jap., Russ., Swed., Swiss and U.S., Leaves, Ger. Herb.; Belg., Dan., Fr. (Jusquame noire) Norw. Port. (Meimandro), Mex. (Beleno Negro), and Span. (Beleno); Leaves and Seeds.

The Brussels Conference agreed to use only the leaf.

Descriptive Notes.—Henbane Leaves occur in commerce in various forms. The dried flowering shoots are known as biennial Henbane, and the first year's large autumnal leaves are sold as annual Henbane. German Henbane consists of the flowering tops of

the small annual form of the plant, which is usually produced from the last formed seeds, and a certain amount of it is usually found in fields of the biennial form. These seeds produce weak plants which flower the first year. It has small leaves, and usually flowers are present, but the drug is less carefully dried than the English plant. Formerly a very superior preparation of the dried flower and leaves of the flowering shoots, with the stalks and mid ribs removed, was sold, but is not at present obtainable. The official drug consists both of fresh leaves and flowers with the branches to which they are attached, and also of the leaves and the flowering tops separated from the branches, collected from the biennial plant, carefully dried. The leaves of the biennial plant are sessile or nearly so, the stalked root leaves decaying as a rule before the plant flowers. The sessile flowers are cup-shaped or irregularly rotate, yellowish with purple veins, and are subtended by large leafy bracts. The fruits open transversely and are two celled, with numerous seeds attached to axile placentæ. The leaves are oblong ovate or triangular ovate, and sinuate in the broad leaved form of the plant and oblong and pinnatifid in the narrow-leaved form. The stout veins and the under surface are furnished with long, clammy, glandular hairs, and the whole plant when fresh has a strong, somewhat unpleasant, but characteristic odour and a slightly acid taste. Recently the leaves and stalks of *Hyoscyamus muticus*, L., have been imported from Egypt. They contain more Hyoscyamine than those of *H. niger*, for which they should therefore not be substituted, but the plant will not flourish in this country. Henbane Leaves in fragments or in powder may be recognised by the small prismatic crystals of Calcium Oxalate (0.010 mm in diameter, *USP*), the 3 to 4 celled hairs with a bicellular or pluricellular gland at the apex, and by the stomata being surrounded by 3 to 4 cells, of which one is smaller than the others.

Tests—*Hyoscyamus* Leaves dried at 100° C (212° F.) contain from 0.06 to 0.15 p.c. of mydriatic alkaloids. The *USP* method of determination resembles that given under *Belladonna Folium*, except that in the place of 10 grammes of the powdered *Belladonna Leaves*, 25 grammes of *Hyoscyamus Leaves* in No. 60 powder are employed, and instead of 50 c.c. of a mixture containing 4 parts by volume of Ether and 1 part by volume of Chloroform, a measured quantity of 100 c.c. of a mixture of similar composition is employed. No method of determination is given in the *P & G*. The ash of the Leaves varies from 8 to 12 p.c. and should not exceed the latter figure.

Preparations.

EXTRACTUM HYOSCYAMI VIRIDÆ. GREEN EXTRACT OF HYOSCYAMUS

A soft Extract, prepared from the juice expressed from fresh Henbane, the albuminous matters being separated at 93.3° C. (200° F.) and rejected.

Dose.—2 to 8 grains = 0.13 to 0.52 gramme

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme.

It is generally used in smaller doses in pills to prevent the griping action of aperients.

Foreign Pharmacopœias—Official in Austri and Belg, alcoholic from dried Leaves, Dan, Norw and Swed, made from Leaves with weak spirit; Dutch and Fr, alcoholic from dried Leaves, Ger and Jap, made with Water and Spirit from fresh Herb, Hung, juice from fresh Leaves, freed from Albumen and evaporated to a thick fluid, equal parts of Spirit added, filtered and again evaporated Ital, from dried Leaves, Alcohol; Mex, from dried Leaves and dilute Alcohol, also Fluid Extract, Port., aqueous from dried Leaves, with Alcohol, Russ, made from Leaves with Water and Spirit, Span, alcoholic from dried Leaves, S, from dried Leaves with dilute Spirit, U S, alcoholic extract from dried Leaves, also Fluid Extract from the same.

The Brussels Conference agreed to prepare a solid extract (containing about 10 p c of Water) by means of Alcohol (70 p c).

Tests—The Green Extract of Hyoscyamus of the B P. is not a standardised preparation. The U S P Extract is required to contain not less than 0.5 p c of mydriatic alkaloids, the Extract, if stronger than this, being diluted with powdered Milk Sugar. The P G Extract is required to yield at least 0.7 p c of alkaloids. The method adopted by the U S P is similar to their method for the determination given under *Extractum Belladonnae* under the exception that, in this instance, 1 gramme of the Extract is employed instead of 10 grammes of the Extract as used in the case of the *Belladonna Extract*. The percentage of mydriatic alkaloids the result of the test is multiplied by 10 instead of by 20.

The P G method of examination is as follows.—A weighed quantity of 2 grammes of the Extract is dissolved in a beaker in 5 grammes of Water and 5 grammes of Absolute Alcohol. 50 grammes of Chloroform are added to this solution and, after a vigorous shaking, 10 c c of a 1 in 3 w/w Sodium Carbonate Solution. The mixture is allowed to stand for one hour, with frequent intervals of vigorous shaking. A weighed quantity of 50 grammes is then filtered through a dry, well-covered filter paper into a flask and about half is distilled. The remaining Ether-Chloroform solution is transferred to a separator. The flask washed with shaken with 10 c c of Hammett's 5 c c of Ether and the combined fluids are added to the separator. The separator is shaken with 10 c c of Hammett's Acid Solution. When the fluids have completely separated sufficient Ether is added to cause the Chloroform-Ether solution to float on the surface of the acid liquid, and the latter is filtered through a filter paper moistened with Water, and the latter is filtered through a 200 c c capacity flask. The Water into a white glass flask of about three successive quantities of 10 c c of Water, the washings being filtered through the same filter, the latter is washed with Water and the combined fluids are added to the separator. The separator is diluted with Water to 100 c c. After the addition of sufficient Ether to form a layer of about 1 cm, 5 drops of Iodoform Solution are added and Hundredth-normal Volumetric Potassium Hydroxide Solution is added until the lower aqueous layer assumes a pale rose coloration, the mixture being shaken after each

addition To produce this colour not more than 6.5 c.c. of the Volumetric Potassium Hydroxide Solution should be necessary

SUCCUS HYOSCYAMI. JUICE OF HYOSCYAMUS

3 of the juice, expressed from fresh Henbane, mixed with 1 of Alcohol (90 p.c.) to preserve it

Dose — $\frac{1}{2}$ to 1 fl. drm. = 18 to 36 c.c.

TINCTURA HYOSCYAMI. TINCTURE OF HYOSCYAMUS

1 of Hyoscyamus Leaves and Flowering Tops in No. 20 powder, percolated with Alcohol (45 p.c.), to yield 10 (1 in 10)

Dose — 30 to 60 minims = 18 to 36 c.c.

Much larger doses, 4 fl. drm. = 112 c.c., have been given in insomnia

Foreign Pharmacopœias — Official in Belg., Dutch, Fr., Span. and U.S., 1 in 10, Port., 1 and 5, also fresh Herb. and Alcohol, equal weights, Mex., 1 in 5 from Leaves, also 1 in 5 from Seeds, also Etheral, 1 in 5. All by weight except U.S.

The Brussels Conference agreed to a strength of 10 p.c., prepared by percolation with Alcohol (70 p.c.)

Tests — Tincture of Hyoscyamus has a sp. gr. of 0.950 to 0.955, contains from about 2.5 p.c. w/v of total solids and about 45 p.c. w/v of Absolute Alcohol. The B.P. is not a standardised preparation. The U.S.P. Tincture is required to contain 0.007 p.c. w/v of mydriatic alkaloids. The method of determination adopted by the U.S.P. is virtually that employed for the assay of the Fluid Extract of Belladonna. A measured quantity of 100 c.c. of the Tincture is evaporated on a water-bath to about one-tenth its volume, sufficient Alcohol (94.9 p.c.) is added to dissolve any separated substance and the resulting liquid is assayed by the process described under Extractum Belladonnæ Liquidum. In calculating the result of the volumetric determination the final multiplication by 10 is unnecessary.

Constant Tincture of Hyoscyamus (Squire) — A Tincture of Hyoscyamus standardised to contain 0.01 p.c. w/v of mydriatic alkaloids, and forming one of the series of Constant Tinctures introduced by Squire in 1888. It has a sp. gr. of about 0.960, contains about 2.5 p.c. w/v of total solids and about 44.5 p.c. w/v of Absolute Alcohol. A sample of B.P. '98 Tincture prepared and assayed in the author's laboratory had a sp. gr. of 0.955, contained 2.64 p.c. w/v of total solids, 44.5 p.c. w/v of Absolute Alcohol, and yielded 0.01 p.c. w/v of mydriatic alkaloids.

HYOSCINÆ HYDROBROMIDUM and HYOSCYAMINÆ SULPHAS. See separate headings

Not Official.

HYOSCYAMI RADIX. — The dried Root of *Hyoscyamus niger* (biennial) collected in the spring. Introduced by Peter Squire in 1878. Contains on the average about 0.15 p.c. of total alkaloid.

Chloroformum Hyoscyami, Linimentum Hyoscyami, and Tinctura Hyoscyami Radicis, are prepared on similar lines to the corresponding preparations of Belladonna.

OLEUM HYOSCYAMI — Hyoscyamus Leaves, 4, Alcohol (90 p.c.), 8; Olive Oil, 40. The leaves are macerated several hours with the Alcohol, then mixed with the Olive Oil and warmed on the water bath till the Alcohol is dissipated. — Ger.

Foreign Pharmacopœias—Official in Austr (Oleum Hyoscyami foliorum coctum), Leaves 100, Alcohol 75, Ammonia 2, Sesame Oil 1000; Belg (Hyoscyami Oleum), Leaves 100, Alcohol 200, Oleum Officinale 1000; also (Hyoscyami Oleum Compositum), Lavender Oil 1, Peppermint Oil 1, Oil of Rosemary 1, Oil of Thyme 1, Hyoscyamus Oil 996 Dutch (Infusum Hyoscyami Oleosum), Leaves 25, Alcohol 50, Ammonia 1, Sesame Oil 250 Fr (Huile de Jusquiame), dried Leaves 1, Alcohol (95 p c) 1, Poppy Oil 10, Jap (Oleum Hyoscyami), Leaves 4, Alcohol 3, Olive Oil 40 Norw and Swed (Oleum Hyoscyami Infusum), Leaves 50, Alcohol 100, Ammonia 1, Sesame Oil 250. Russ (Oleum Hyoscyami), dried Leaves 4, Alcohol (90 p c) 3, Sesame Oil 24 Span (Aceite de Beleno), fresh Leaves 5, Olive Oil 10 Swiss (Oleum Hyoscyami), Leaves 10, Alcohol 10, Ammonia 2, Sesame Oil 100, also (Oleum Hyoscyami Compositum) (*Syn* Balsamum Tranquilli) same as Belg

The majority of the above work out about 1 of Leaves in 10 of product

HUILE DE JUSQUIAME COMPOSÉE (Baume Tranquille. (Fr) — Dried Leaves of Belladonna, Henbane, Black Nightshade, Poppy and Stramonium, of each 5, Oils of Lavender, Peppermint, Rosemary and Thyme, of each 1, Alcohol (95 p c), 200, Poppy Oil, 5000

Moisten the powdered leaves with the Alcohol, and digest on a water-bath for 24 hours, add the Poppy Oil and heat for 6 hours at 60° to 70° C, stirring occasionally, express, allow to settle, and decant, add the Oils and filter.

Oleum Hyoscyami Infusum — Hyoscyamus — Alcohol (95 p c), 15, Ammonia Water (U S P), 0 4, Oil, 50

Moisten the powder with Alcohol and Ammonia previously mixed, pack tightly and cover well, and macerate for 24 hours, add 12 of mixed oils, digest with agitation for 12 hours at a temperature between 50° and 60° C, strain and express To the residue add the remainder of the Oils, digest and express as before and mix the expressed portions — U S N F

This process is a modification of that prescribed by the Codex and may be used for similar Infused Oils

Oleum Hyoscyami Compositum *Syn* Balsamum Tranquillans — Oils of Absinth, Lavender, Rosemary, Sage, Thyme, of each 2 drops, Infused Oil of Hyoscyamus (N F) 100 cc — U S N F

HYOSCYAMINA Hyoscyamine $C_{17}H_{23}NO_3$, eq 287.05 — A crystalline alkaloid obtained from the Seeds of *Hyoscyamus niger*, the Root of *Scopolia carniolica*, and probably other allied plants, isomeric with Atropine but not identical with it

It occurs as white needle-shaped crystals Only slightly soluble in Water, but freely in Alcohol (90 p c), in Chloroform, and in Ether Probably constitutes the greater portion of the crystallisable alkaloid naturally existing in all the mydriatic drugs, and best obtained from the Root of *Scopolia* or *Belladonna* Most of the commercial 'Atropine' consists principally of Hyoscyamine

The salts used in medicine are the Hydrobromide and Sulphate

Dose — $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0 005 to 0 001 gramme

Hager, maximum single dose, 0 005 gramme, maximum daily gramme

Tests.—Hyoscyamine melts at 108.5° C (227.3° F) Its neutral solvents are alkaline in reaction towards Litmus Solution and leaves a white residue It forms with Auric Chloride a Gold double salt melting at 162° C (320° to 323.6° F) It dissolves in Sulphuric Acid without change of colour, and no alteration in colour should occur on the addition of one or two drops of Nitric Acid It should leave no residue when ignited with free access of air

HYOSCINA Scopolamine, Hyosine $C_{17}H_{21}NO_3$, eq. 318.81 — An alkaloid which is found in *Hyoscyamus niger*, and various species of *Scopolia*. It now represents what was formerly used in medicine under the name 'Amorphous Hyoscyamine' It is usually employed in the form of Hydrobromide, Hydrochloride, and Hyosine

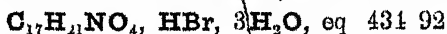
Foreign Pharmacopœias—Official in Mex

It forms transparent moderate sized crystals or a colourless transparent glassy mass. It is slightly soluble in Water, readily soluble in Alcohol (90 p c), Ether, Chloroform and diluted acids.

Tests—Crystalline Hyoscyne, when dry, melts at 50°C (122°F). When dried over Sulphuric Acid the crystals lose in weight and change to a colourless amorphous glassy looking mass which will not recrystallise. Its faintly acidified solution gives with Potassium mercuric Iodide (Mayer's) Solution a yellowish-white precipitate, with Mercuric Chloride Solution a white precipitate, with Picric Acid a yellow crystalline precipitate. Tannic Acid produces no precipitate. Auric Chloride Solution added to a solution of Hyoscyne faintly acidified with Hydrochloric Acid yields a yellow precipitate which, recrystallised from Water, yields brilliant yellow, glistening needles, which melt at 212° to 214°C (413° to 417°F). It leaves no weighable residue when ignited with free access of air.

HYOSCINÆ HYDROBROMIDUM.**HYOSCYNE HYDROBROMIDE.**

B.P. Syn—HYDROBROMATE OF HYOSCYNE, SCOPOLAMINE HYDROBROMIDE.



Fr., BROMHYDRATE D'HYOSCINE, CH. R., SCOPALAMINHYDROBROMID,
Ital., BROMIDRATO DE SCOPOLAMINA

Colourless, transparent, rhombic crystals, permanent in the air.

A similar description is common to the *B.P.* and *U.S.P.*

It is the Hydrobromide of an alkaloid Hyoscyne (Scopolamine) obtained from Hyoscyamus, various species of Scopolia and other plants of the Solanaceæ.

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from the light.

Atropine, the crystalline variety of Hyoscyne, forms a crystalline Hydrobromide.

Solubility—1 in 4 of Water, 1 in 14 of Alcohol (90 p c); very slightly soluble in Chloroform or Ether.

B.P. states it is 'soluble in one part of cold Water,' which is incorrect, 1 in 4 is more nearly so.

Medicinal Properties—Hypnotic and sedative. Highly recommended in all forms of violent mania and cerebral excitement.

Given by the mouth it appeared (*M.P.* '05, i 615) to be of great benefit in acute mania, giving quiet sleep, whereas by hypodermic injection it caused an alarming degree of depression.

In paralysis agitans (*B.M.J.* '05, ii 12), 0.2 to 0.3 mgr, either in pill or in solution. In certain cases of senile pruritus it was of value in doses of 0.8 to 0.5 mgr daily. In spasmodic asthma it was administered in large doses (0.25 to 0.5 centigramme) subcutaneously with Caffeine, with very great relief. In acute mania hypodermic injection resulted in eight or nine hours' sleep, with consequent improvement in the general mental state.

The racemic form seemed (*B.M.J.* '05, ii 250) to be less liable to produce untoward effects as a hypnotic, and was of equal hypnotic value, but Hyoscyamine and Hyoscyne should be used with caution.

Exists in two forms (*B.M.J.* '05, ii 1005), one of which is optically active (laevorotatory), whilst the other is indifferent to the ray of polarised light or racemic. The two have the same effect as hypnotics, but the racemic form has only half the action of the laevorotatory base on the pupil, glands and heart.

Scopolamine the best of all sedatives in the vomiting of pregnancy—*B.M.J.* '07, ii 27.

1000 confinements conducted with its assistance — *B M J W.* '07, n. 10

In epileptic attacks of an hysterical form — *M J* '95, 244

As a mydriatic (1 grain to 1 oz), in cases where Atropine is undesirable — *B M J* '94, n. 598

Incipient acute mania arrested by a single injection of $\frac{1}{100}$ grain — *B M J.* '97, n. 652

In mania, $\frac{1}{100}$ grain hypodermically and after forty minutes $\frac{1}{100}$ grain by the mouth to procure seven hours' sleep, followed after an interval of two days by a dose of $\frac{1}{8}$ grain hypodermically to induce ten hours' sleep. Sodium Bromide in firm doses being administered during the interval and for two days after the second sleep until 2 oz in all had been taken — *B M J* '08, 1 71

In the palliative treatment of paralysis agitans, it is probably the most useful drug that has hitherto been tried, $\frac{1}{100}$ to $\frac{1}{80}$ grain in solution in Chloroform Water, administered by the mouth (great caution required as regards the dose, and it is well not to begin with more than $\frac{1}{100}$ or $\frac{1}{80}$ grain. $\frac{1}{4}$ grain given two or three times a day (by the mouth), for long periods without noting any bad effects — *P. M.* 410

In exophthalmic goitre, $\frac{1}{100}$ grain — *M J* '02, 280

Two cases of paralysis agitans treated with Hydrobromide, at first hypodermically, the dose being gradually increased from $\frac{1}{100}$ to $\frac{1}{80}$ grain, injected once a day. Subsequently administered in $\frac{1}{80}$ grain doses dissolved in Chloroform Water, given twice daily by the mouth, gradually increasing the dose up to $\frac{1}{4}$ grain — *L* '02, 1 1907

Hyoscine is more sedative and more reliable as a hypnotic than Hyoscyamine. Indeed, it has almost totally replaced it for this purpose. There can be little doubt, moreover, notwithstanding the many ill effects attributed to the use of Hyoscine (Scopolamine), and to its variable action, that as a hypnotic it has come to stay. Its solubility in Water and its applicability to hypodermic medication make it of extreme value in many conditions, particularly in the insane — *L* '99, n. 142

$\frac{1}{100}$ grain, and subsequently $\frac{1}{80}$ grain every thirty minutes to one hour, for from twenty-four to forty-eight hours, until the patient has taken from forty to sixty doses, in the treatment of the drug habit — *T G* '02, 41, 71

Morphine-Scopolamine Anaesthesia — Hypodermic injection of $\frac{1}{100}$ to $\frac{1}{80}$ or even $\frac{1}{4}$ grain of Scopolamine Hydrobromate with $\frac{1}{4}$ grain of Morphine, to be repeated after one or two hours, previous to an operation. Very little Chloroform is required, and the patient's sleep for hours after the operation, and do not complain of any pain — *L* '12, 1 44

Morphine-Scopolamine anaesthesia would be found useful in those cases where Chloroform and Ether are both contra-indicated, but that its action is not narcotic enough to admit of its taking the place of the general inhalation anaesthetics — *B M J E* '03, 1 14

G. Volkmann found it advisable to give 12 milligrammes (about $\frac{1}{4}$ grain) of Scopolamine and 15 centigrammes (about $\frac{1}{100}$ grain) of Morphine four hours before the operation and ten centigrammes (about $\frac{1}{100}$ grain) two hours later, and $\frac{1}{4}$ hour before the operation he gives 9 mg (about $\frac{1}{100}$ grain) of Scopolamine and 5 mg (about $\frac{1}{100}$ grain) of Morphine. In the case of old people, or patients suffering from diseases of the internal organs he has employed smaller doses. In some cases the anaesthesia had to be deepened by Ether inhalation (given drop by drop) — *B M J E* '04, 1 21

Administered hypodermically in repeated doses of $\frac{1}{4}$ grain accompanied by $\frac{1}{4}$ grain Morphine in the afternoon, four hours, two hours, and one hour before the time fixed for operation, it produces (M J '05, 1. 575) a quite restful sleep, giving two or three hours good anaesthesia. It does not however, give complete muscular relaxation, and the patient ought not to be given in operations where complete relaxation is required, e.g., abdominal operations.

As a general anaesthetic, a solution of 1 milligramme ($\frac{1}{100}$ grain) of Scopolamine in 1 c.c. of Distilled Water injected subcutaneously four hours before the surgical operation, followed by a second injection after an interval of one hour (*B M J* '06, 1 100)

Scopolamine-morphine is used as a preoperative anaesthetic, as a preparative to Chloroform.

this is supported by Korff, Kümmel, Krönig, and others. For this purpose, small doses, $\frac{1}{2}$ milligramme ($\frac{1}{120}$ grain) of Scopolamine with $\frac{1}{2}$ to $1\frac{1}{2}$ centigrammes ($\frac{1}{4}$ to $\frac{1}{2}$ grain) of Morphine, should be given half to one hour before the operation, or better still, if time allows, half this dose should be given an hour and a half before operation, and repeated in an hour. Chloroform must be given in very small quantities, drop by drop, and should not be pushed to the point of abolishing the reflexes. This persistence of the reflexes, indeed, is one of the disadvantages, and the limbs should be mechanically controlled. Ether is still less satisfactory in this respect. In old people, however, even the small dose of Scopolamine is sometimes sufficient to induce anaesthesia without the addition of Chloroform. At the end of the operation, the patient should be given an infusion or enema of 1 to 2 pints of warm Saline Solution, in order to obviate the thirst which in these cases is sometimes distressing, and when he awakes some hours after he may be given a little food. Special care must be taken with children and patients suffering from diseases of the heart or kidneys. See also *B M J* '05, ii 185, and *B M J* '07, i 47.

Two cases in which the subcutaneous injection of 1 mg, along with 1 cg. of Morphine, one hour before the administration of chloroform was followed by death on the operation table—*B M J* '07, ii 68.

Dose— $\frac{1}{200}$ to $\frac{1}{100}$ grain = 0.0003 to 0.0006 gramme

Ph. Ger. maximum single dose, 0.001 gramme, maximum daily dose, 0.008 gramme

Prescribing Notes—Best given by hypodermic injection. When given by the mouth it may be conveniently dissolved in Chloroform Water.

Not Official—Guttæ Hyoscinæ, Guttæ Hyoscinæ et Cocainæ, Hyoscine Dises, Injectio Hyoscinæ Hypodermica, Hyoscinæ Hydrochloridum (Scopolamine Hydrochloride), and Hyoscinæ Hydriodidum (Scopolamine Hydriodide).

Antidotes—Pilocarpine Nitrate, half a grain hypodermically, or $\frac{1}{4}$ grain Morphine, then stomach-tube or emetic, followed by stimulants and artificial respiration.

Foreign Pharmacopœias—Official in Dan., Dutch, Ger., Jap., Swiss and U.S. (Scopolaminum Hydrobromidum), Ital. (Bromhidrato di Scopolamina). Not in the others. The title 'Hyoscine Hydrobromide,' introduced into *Ph. Ger.* iii, has been replaced in *Ph. Ger.* iv by 'Scopolamine Hydrobromide.'

Tests.—Hyoscine Hydrobromide contains theoretically 12.38 p.c. of Water. It is officially required to lose rather more than 12.0 p.c. of its weight at a temperature of 100° C (212° F), the *P.G.* gives 12.3 p.c., the *U.S.P.* states that it loses its Water of crystallisation at 110° C (230° F). The anhydrous salt melts, according to Hesse, at 181° C (357.8° F), and not, as officially stated at 193° to 194° C (379.4° to 381.2° F). Jowett confirms Hesse's melting point, and states that the purified lævo salt melts at 193° C (379.4° F), and the inactive modification at 180° C (356° F). The tests and characters of the official salt should therefore be given for the pure product as it appears in commerce, which is a mixture of stereoisomers melting at 181° C (357.8° F). The *U.S.P.* gives the melting point as 191° to 192° C (375.8° to 377.6° F), the *P.G.* at about 180° C (356° F). Its aqueous solution is stated in all three Pharmacopœias to be slightly acid in reaction towards Litmus, though Jowett states (*P.J.* '98, ii 196) that there is no reason why the salt should not be neutral to Litmus. Its aqueous solution slightly acidified with Hydrochloric Acid yields with Potassiummercuric Iodide (Mayer's) Solution, a yellowish-white precipitate.

Its aqueous solution yields with Mercuric Chloride T.S. a white precipitate, with Phospho-tungstic Acid Solution a white precipitate, when in sufficiently concentrated solution it yields with Picric Acid a yellow precipitate, with Iodine Solution a brown precipitate, and with Platinum Chloride Solution a white precipitate. The aqueous solution yields a whitish precipitate with Potassium Hydroxide Solution. The turbidity is only produced on the addition of a considerable excess of Sodium Hydroxide Solution and disappears quickly. The aqueous solution is not precipitated by Ammonia Solution, or by Potassium Bichromate Solution. It is officially stated to form a crystalline salt with Auric Chloride having a m.p. of 198°C (388.4°F); Jowett (*JCS Trans*, '97, 679) has shown that under the B.P. conditions an additive compound Hyoscine Hydrobromide Gold Chloride melting at 215°C (419°F) is formed, but that when prepared in the usual manner the Aurichloride melts sharply at 198°C (388.4°F). The U.S.P. gives the m.p. of the pure Chloraurate at 197°C (386.6°F). When Hyoscine Hydrobromide is dissolved in Water it yields with Silver Nitrate Solution a yellowish curdy precipitate, insoluble in Nitric Acid, and when washed, practically insoluble in Ammonia Solution. One or two drops of Chlorine Water added to a small quantity of a 1 in 10 aqueous solution yield a reddish-brown solution, and when shaken with Chloroform the brownish-red colour passes into the chloroformic layer. A small crystal of the salt evaporated to dryness in a white porcelain dish on a water-bath leaves a yellowish residue, which upon the addition of a few drops of Potassium Hydroxide Solution yields a violet coloration. Its freedom from readily charred organic impurities may be ascertained by the Sulphuric Acid test, the salt should yield but a pale yellow coloration when treated with this Acid, and if after the addition of Nitric Acid no colour is developed the absence of Morphine may be inferred. The salt should leave no weighable residue when ignited with free access of air.

Commercial samples may contain in addition to inactive Scopolamine (Atropine) Apotropine. Apotropine is, according to Kobert (*PJ* '05, i 442), a dangerous impurity and one to be rigidly excluded. Absolutely necessary for the official description to require a rotatory power not less than that shown by the pure salt -25.45° for a 6.5 p.c. solution at 15.8°C (60.5°F).

Sulphuric Acid—Only a faint yellow colour should be developed on the addition of Sulphuric Acid to Hyoscine Hydrobromide.

Nitric Acid—On the subsequent addition of a drop of Nitric Acid to the above mixture no coloration should be developed, U.S.P.

Not Official.

GUTTÆ HYOSCINÆ (*Syn* Guttæ Scopolaminæ.—Hyoscine Hydrobromide, 2 grains, Distilled Water, 1 fl oz.—*London Ophthalmic and Charing Cross*

Hyoscine Hydrobromide 0.5 or 1 p.c.—*St Thomas's and B.P.C.*

GUTTÆ HYOSCINÆ ET COCAINÆ—Hyoscine Hydrobromide, 0.5 p.c., Cocaine Hydrochloride, 1.0 p.c.—*St Thomas's and B.P.C.*

INJECTIO HYOSCINÆ HYPODERMICA—For parenteral use is made by dissolving Hyoscine Hydrobromide, 1 grain, in distilled Water, 500 minims, but the strength should always be stated.

Dose—2 to 5 minims = 0.12 to 0.3 c.c. as a sedative in nervous diseases, especially where there is much violence and excitement. When given by the mouth at least double the dose is required to produce the same effect.—*L.* '89, ii 736

Hyoscine Discs— $\frac{1}{100}$ and $\frac{1}{75}$ grain, *St. Bartholomew's*, $\frac{1}{100}$ grain, *Guy's*

HYOSCINÆ HYDROCHLORIDUM (Hyoscine Hydrochloride, Scopolamine Hydrochloride).—Large, colourless, prismatic crystals, or as a colourless crystalline powder, readily soluble in Water, and in Alcohol (90 p.c.)

Dose— $\frac{1}{100}$ to $\frac{1}{60}$ grain = 0.0003 to 0.0006 gramme

Tests—Hyoscine Hydrochloride answers to the tests distinctive of Hyoscine given under Hyoscine and Hyoscine Hydrobromidum. The aqueous solution yields with Silver Nitrate Solution a white curdy precipitate, which, when filtered and washed, is insoluble in Nitric Acid but readily dissolves in Ammonia Solution. The salt when ignited with free access of air leaves no weighable residue.

HYOSCINÆ HYDRIDIDUM (Hyoscine Hydriodide, Scopolamine Hydriodide).—Colourless, transparent prismatic crystals. Soluble in Water, and in Alcohol (90 p.c.)

Dose— $\frac{1}{60}$ to $\frac{1}{100}$ grain = 0.0003 to 0.0006 gramme

HYOSCYAMINÆ SULPHAS.

HYOSCYAMINÆ SULPHATE

($C_{17}H_{23}NO_3$), H_2SO_4 , $2H_2O$, eq. 707.20

White, slender, crystalline, hygroscopic needles, or an odourless, white, granular, hygroscopic powder.

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with air, especially moist air.

Solubility—2 in 1 of Water, 1 in $4\frac{1}{2}$ of Alcohol (90 p.c.), very slightly soluble in Chloroform or Ether.

Medicinal Properties.—In small doses it is a sedative for mental excitement and insomnia, and in large doses it has been used for calming the excitement of delirium tremens and acute mania, but for this purpose it is superseded by the salts of Hyoscine.

Taken for sea-sickness in $\frac{1}{60}$ grain doses three or four times a day, two or three days before embarkation, and for the first days on board ship, until nausea has disappeared.—*B. M. J.* '91, ii 576.

These are conveniently carried as half-grain pilules, made with 'Diluting Mixture,' p. 897.

Dose. $\frac{1}{60}$ to $\frac{1}{100}$ grain = 0.0003 to 0.0006 gramme

Haqer, maximum single dose, 0.005 gramme, maximum daily dose, 0.015 gramme.

Not Official. Hyoscyaminæ Hydrobromidum, Hyoscyaminæ Discs

Foreign Pharmacopœias.—Official in U.S. Not the others.

Tests. Pure Hyoscyamine Sulphate melts at $204^{\circ}C$ ($399.2^{\circ}F$), the commercial Hyoscyamine Sulphate melts at about $200^{\circ}C$ ($392^{\circ}F$), the official melting point is $206^{\circ}C$ ($402.8^{\circ}F$). The U.S.P. m.p. is $198.9^{\circ}C$ ($390.1^{\circ}F$). Jowett suggests (*P. J.* '98, ii. 196) that an official m.p. should be given not lower than $200^{\circ}C$. ($392^{\circ}F$). Its aqueous solution is neutral in reaction.

towards Litmus paper and is levogyrate. It affords a yellow precipitate with Auric Chloride. Both the *BP* and *U.S.P.* state that the yellow precipitate is soluble in boiling Water acidified with Hydrochloric Acid, and again deposited in the form of brilliant golden yellow scales on cooling the solution. A small crystal evaporated to dryness in a porcelain dish on a water-bath, with 5 drops of Nitric Acid, leaves a yellowish residue which, when moistened with Alcoholic Potassium Hydroxide Solution, affords a purple-violet coloration. The m.p. of the Aurichloride is 160°C . (320°F). The *U.S.P.* gives this figure for the m.p., but *BP* gives no figure. The *U.S.P.* also gives a figure for m.p. of the Picric Acid. The aqueous solution of the salt yields with Barium Chloride Solution, a white precipitate insoluble in Hydrochloric Acid.

The more generally occurring in Hyoscine impurities are fixed residue, alkaloids other than Hyoscyamine, readily charred organic impurities.

The salt should leave no residue when ignited with free access of air. Most alkaloids other than Hyoscyamine may be detected by the behaviour with Potassium Chloride Solution as described in the small type below, charred organic impurities are revealed by the Sulphuric Acid test. Hyoscyamine may be distinguished from Atropine by its optical activity and by the test of its Aurichloride. Compare Atropine, p 199.

Picric Acid—Hyoscyamine Picric Acid melts at 175°C (347°F), *U.S.P.* Atropine Picric Acid melts at 162°C (324°F), *BP*.

Platinic Chloride—No precipitate is formed in solutions of the salt on the addition of T.S. of Platinic Chloride, *U.S.P.*, a solution in Water acidulated with Hydrochloric Acid yields no precipitate with Platinic Chloride Solution, *BP*.

Sulphuric Acid—No colour could be produced when Sulphuric Acid is added to Hyoscyamine Sulphate, *U.S.P.*

Not an Official

HYOSCYAMINÆ HYDROBROMIDE—Short white or yellowish-white prismatic crystals, readily soluble in Water, and in Alcohol (90 p.c.). On account of its hygroscopic nature it should be kept in well-stoppered glass bottles of a dark color, and protected as far as possible from exposure to air.

Dose— $\frac{1}{100}$ to $\frac{1}{10}$ grain = 0.002 to 0.006 gramme

Official in U.S.

Tests—Hyoscyamine Hydrobromide melts at 151.8°C (305.3°F). Its aqueous solution is neutral in reaction towards Litmus and is levogyrate. It answers the tests distinctive of Hyoscyamine given under Hyoscyamine Sulphate, with the exception of the test with Gold Chloride Solution. The *U.S.P.*, 1890 stated that with Gold Chloride Solution an aqueous solution of the salt yields a precipitate which when recrystallised from a small quantity of boiling Water acidulated with Hydrochloric Acid, is deposited on cooling in minute lustrous golden yellow scales.

Jowett has shown (*J.C.S. Trans.*, '97, 679) that under the above conditions Gold Chloride is produced, it forms a yellowish-red precipitate on cooling in acidified Water in scales, melting sharply at 164°C (327.2°F) and suggesting that the *U.S.P.* description required modification. The description of the test first appearing in the 8th Decennial Revision of the *U.S.P.* is, however, the same as that given in the 1890 Edition. The test solution yields with Silver Nitrate Solution a white precipitate which is insoluble in Nitric Acid, while

Solids by Weight

insoluble in Ammonia Solution, but reliable to contain the same impurities for their detection in the latter may be test for Morphine, no colour should be to a mixture of Hyoscyamine Hydrobromide.

Hyoscyamine Dises (for hyp) also applied here. The USP includes a Hyoscyamine Sulphate — *St Bartholom* produced on the addition of Nitric Acid and Sulphuric Acid

dermic injection) - $\frac{1}{8}$ and $\frac{1}{2}$ grain of
Not C.W.'s

Not Clw's

ICHTHY —

being official

FR, CORRE DI POISSON, CIR, HAL
SPAN, IC OCOLLA.

The swimming bladder or sound of vu^{CLASS}
ent into two sheds. SPENBLAST, ITAL, COLLIER DE PESCARI;

This well known substance was in the Pharmacopoeia called Ichthyocola or Fish Glue, it was still to be found in most of the Continental Pharmacopoeias, and was used in the early London Pharmacopoeias, and the best quality Isinglass is used for Cod Liver Oil, and is used in medicine as a nutrient. It is

This is included among the Tests of Tannin Acid, with which it forms an insoluble compound. It is used in medicine as a nutrient. It is used for finishing Yellow Leather, and is also used in the manufacture of Gunpowder. Russian Ienglass is reckoned the best quality. Ienglass is also used in the manufacture of Gunpowder.

Foreign Pharmacopœias Official in
Norw. and Russ. (Colla Piscium), US
de Pescadi), Port. (Gelatina de Peix
others

Descriptive Notes—There are many varieties of Isinglass in commerce differing in shape and quality. It consists of the swimming bladder of various fishes washed and freed more or less from the membrane. The kind prepared for use in medicine is derived from the bladder of the *Huso*, etc) and is imported from Russia. The Isinglass, as the swimming bladder is called, is twisted when soft into various shapes or forms, such as threads, which can be distinguished from Gelatin by its transparency and its melting point. It is prepared in the form of Isinglass threads, which can be distinguished from Gelatin by its transparency and its melting point. It is prepared in the form of Isinglass threads, which can be distinguished from Gelatin by its transparency and its melting point.

Tests—Isinglass is not soluble in cold Water, but is dissolved and known as long and short almost entirely in boiling Water. On treating here is offered cut up into slender the substance swells uniformly, producing a white gelatin by its laminated structure dissolves entirely, (gelatin, under similar conditions, but the best qualities dissolve a nearly transparent solution. The best Russian Isinglass with hot Water, in which ordaine jelly which gradually

Not Official.

ICHTHYOL^{me}

AMMONIUM RUTHYOSUL.

A reddish brown, syrupy liquid, with ignoi
Obtained by the action of Sulphuric Acid on a
distilled from peculiar fossil deposits, principally
tion with Ammonia

Solubility.—Entirely soluble in Water, partially in Alcohol, and Ether, entirely in a mixture of both.

It mixes readily with Glycerin, Fats, Oils, Soap

The following formula is recommended—*L* '33, 1. 354 It is better to boil Acetic Acid, 30, boil down to 20, add Olive Oil, 10, all by weight, to make an Ointment. If evaporated only to 20 as down to 13, as Water separates from the Glycerin as a tampon—*L* '90, 1. 1142, directed

For uterine affections it is used with '94, 11 1113 As a paint (20 p.c. sol.)
'91, 1 55 antiseptic injection in vesical catarrh,
pharyngitis — 73 349 P. 11 370 1 to 2 p.c. aqueous

'91, 1 55 As a gargle in acute pharyngitis—*L* '83, 349, *P* '81 370 1 to 2 p.c. aqueous for foot blisters—*T G* '93, 56 As 10 p.c.—*L* '97, 1 1165, *T G* '96, 350 *M A* '95, 139, and in gonorrhoea *T G* '91 sometimes beneficial in some hyper-solution used as irrigation in gonorrhoea. 1 785

Given to a limited extent and has beryl, 1, Vaseline, 1, applied to the pus-
-emic diseases like acne rosacea.—L '08, 1—B M J E '08, 11 24
—T G '99, 819.

As an ointment composed of lenthil made with Lard — *I G* '93, 813.

In prairies, vultures, as a 15 p c ointn 95, n 28, P J '95, n 51, '96, n 494,
In 3 prairie doves in urticaria — B N

In 3-grain doses in urticaria — *B M J E* '94, 1 1521, *B M J E* '95, 1 51, skin beyond the affected part modifies, 'f, erysipelas, 30 to 60 pc Ointment, *B M J E* '99, 1 60

Applied so as to cover the healthy skin — *TG* '91, 862, '92, 294, 684, *M.I.*
and distinctly shortens the duration of 5 to 10 grain suppository in prostatitis.

95, 249, *B M J E* '94, 1 24, 48, as application of 1 Ichthyol and 5 Vaseline to
— *B M J E* '93, 11 24, for twelve hours, but it completely

It is not without danger, as an af-
a child for ... produced stu-
recovered—B I J ... 1018
... Oleatis, 10, Aq Calcis, 10, Ichthyol,

Zinc Oxide, 20, Magnes Carb, 10, 95, 11 92
the first degree Calcu Carb, 10, 2 grammes
1 to 3, for extensive burns — *B M J E*
made with a mixture of *Althaea* 3, *Inquice*

Dose—15 to 30 grains = 1 to 2 g. **Preparations**—In pill form. Also given in capsules.

Powder 3, and Compound Tragacanth
num Ichthyol or to 4 of Sodium Ich
pressed Tablet.

The Oils of Citronella, Turpentine, and Bergamot. Essence of Almonds is also for disguising the odour of Linseed Oil. Milk, Chocolate, or Oil of Peppermint. Ammonia in salt is generally the best for this purpose. It is more or a solid, and makes a "r" sound.

When Ichthyol is ordered, the Sodium salt has similar properties. Combined with Vaseline or Lanolin it forms a 10 to 20 p.c. Ointment, also a 10 to 20 p.c. Collodion. Zinc Ichthyolsulphonates have been employed as a 10 to 20 p.c. Ointment, also a 10 to 20 p.c. Collodion.

Lithium, Magnesium and Ammonium salt makes a suitable pill-mass, and may be prepared by making 120 grains of the Ammonium salt and 15 grains of light Calced Magnesia into a pasty or two of Water.

This orange powder Bromide, a turbid brown precipitate settling salt, will make nice pills with a & adheres to the bottle, the addition of Mucilage When dispensed with Potas

to a sticky mass & thrown out, as with Ichthyol after a time becomes hard and of Acacia does not prevent this. made with Cocoa-butter alone 3 grains of For pessaries a Gelatin Theobroma make a good suppository

Incompatibles —Alcohol, Potassium Bromide. All sulphate, and decompose.

alkaloid, and liberation of Ammonia With alkaloidal salts a double decomposition takes place.

Foreign Pharmacopœias—Belg, Ital (Ichthol), Jap, Ammonium Sulphoichthyolate, Russ and Span (Ictiol)

Tests—Ammonium Ichthiolsulphonate, when warmed with Potassium or Sodium Hydroxide Solution, evolves Ammonia gas, readily recognised by its odour and by its action upon moistened red Litmus paper, if the mixture be evaporated to dryness and ignited a carbonaceous mass is left, which evolves an odour of Hydrogen Sulphide when oxidised with Hydrochloric Acid When evaporated on a water bath it usually loses about 15 p.c. of its weight, and should lose at the most not more than 50 p.c. The clear aqueous solution is slightly alkaline in reaction towards red Litmus paper A 10 p.c. aqueous solution, when mixed with Hydrochloric Acid, throws down a dark resinous precipitate, which is soluble in Ether and in Water, but is reprecipitated from the latter liquid by Hydrochloric Acid or Sodium Chloride When evaporated and ignited with free access of air it should leave no weighable residue

GELATUM ICHTHYOL Gelatin, 1, Distilled Water, 2½, Ichthyl, 1; Glycerin, 6 all by weight—*Pharm. Form.*

This has been incorporated in the *B.P.C.* under the title *Pasta Ichthamolis*.

INJECTIO ICHTHYOL 2 to 5 p.c.—*Loch*

PASTA ICHTHYOL (Unit) Ammonium Ichthyolate, 2 scruples to 2 drms., Powdered Dextrin 1 oz. Distilled Water, 1 oz., Glycerin, 6 drms. Dissolve the Ichthyl in the Water and Glycerin, mix with the Dextrin and heat on a water bath until uniform *Pharm. Form.*

Ammonium Ichthyl 2½, Carbolic Acid 2½ Dissolve in warm Water 22½ and Starch 50—*B.M.J.* '91, i 101

This has been incorporated in the *B.P.C.* under the title *Pasta Ichthamolis Composita*.

Dr. Unna considers that for certain purposes a waterless Ichthyl varnish possesses advantages over the usual preparations, and gives the following formula Ichthyl, 40, Starch, 40 Solution of Albumen, 1 to 1½, Water, to 100 The Starch is first moistened with the Water then the Ichthyl well rubbed up with it, and lastly the solution of Albumen is added—*L.* '91, i 622, *B.M.J.E.* '91, i 102

UNGUENTUM ICHTHYOL Ichthyl 1, Paraffin Ointment, 9, Mix.—*King's*

Recommended in the treatment of chaldam.—*B.M.J.* '91, i 503

Unguentum Ichthamolis—Ammonium Ichthyosulphonate, 10, Hydrous Wool Fat, 90, Mix *B.P.C.*

UNGUENTUM ICHTHYOLIS—Ichthyl, 40 grains; Salicylic Acid, 8 grains, Soft Paraffin, to 1 oz. *London*

UNGUENTUM ICHTHYOLIS COMPOSITUM—Ichthyl, 1, Solution of Lime, 9, Hydrous Wool Fat, 5, Soft Paraffin (yellow), 10, Zinc Ointment, 5.—*Guy's*

VASOLIMENTUM ICHTHYOL—Ammonium Ichthyl, 10, Liquid Vasoliment, 90—*Haq.*

Parogen Ichthamolis See Ichthamolis Vasoliment—Ammonium Ichthyosulphonate, 10, Parogen, 90 to produce 100—*B.P.C.*

NATRIUM SULPHO-ICHTHYOLICUM—Sodium Ichthyolsulphonate).—A brownish black tar like mass, with a bituminous odour

Solubility. It makes a somewhat turbid solution with Water; dissolves in a mixture of equal weights of Alcohol and Ether. It is soluble in Benzol.

Medicinal Properties. The same as the Ammonium salt

Tests.—Sodium Ichthyl dissolves in Water, forming a solution which is only faintly alkaline in reaction towards red Litmus paper When ignited it

leaves a residue possessing an alkaline reaction which colours a non-luminous flame intensely yellow, and which, when dissolved in Water and acidified with diluted Nitric Acid yields, with Barium Chloride Solution, a white precipitate insoluble in Hydrochloric Acid. The aqueous solution, when mixed with Hydrochloric Acid, precipitates a dense cream-coloured mass which, when separated from the supernatant liquid, is soluble in Ether and in Water, but is again precipitated from the latter fluid by the addition of Hydrochloric Acid or Sodium Chloride. It contains from 25 to 30 p.c. of moisture, which may be determined by drying over Sulphuric Acid. It is not a true desiccator. It should not evolve Ammonia when treated with Sodium Hydroxide Solution, indicating the absence of Ammonium Ichthyosulphonate.

ICHTHOFORM (Formaldehyde Ichthyosulphonate)—A blackish-brown powder, possessing a strong odour, insoluble in Water, and in Alcohol (90 p.c.). Introduced in 1891 as a disinfectant.

Dose—10 to 20 grains = 0.65 to 1.3 grammes

FERRICHTHOL (Iron Trichthosulphonate)—A dark, blackish brown, non-hygroscopic, amorphous powder. Has been given in anaemia.

ANYTIN—Under this title a 33 p.c. solution of Ichthyosulphonic Acid has been introduced into medicine. Its property of rendering soluble in Water substances which are otherwise insoluble is so marked that the compounds so produced are known as Anytols, as Mera-Cresol Guaiacol, Camphor and Iodine-Anytols have received some attention as antiseptic agents, chiefly as antiseptics.

THIOL—An artificial substitute for Ichthyol, imitating the action of Sulphur on gas oil, and subsequent treatment with Nitric Acid. It is supplied in two forms, a powder and a liquid, it is soluble in Water and almost odourless.

Useful in acute forms of erysipelas, in erythema, and in inflammatory diseases of women, also in pruritus of the rectum generally. *Pharm.* 565

A 20 to 40 p.c. solution is used for erysipelas in the manner as Ichthyol. — *B.W.J.F.* '94 103, *L.C.* '94, 627

ICHTHALBIN (Albumen Ichthyosulphonate)—A greyish-brown powder, almost odourless and tasteless. Insoluble in Water, decomposed by heat.

Dose— $7\frac{1}{2}$ to 30 grains = 0.5 to 2 grammes per diem

Tumenol—A similar body to Ichthyol, is a thick dark brown liquid. It is a mixture of Tumenolsulphonate (Tumenol Oil) and Tumenolsulphonic Acid (Tumenol Powder).

Tumenol Ammonium is a compound introduced to overcome the difficulty with which ordinary Tumenol is miscible with various diluents. It contains more Water than Tumenol. It is practically neutral and is soluble in Water, is miscible with slight turbidity to the extent of 1 to 5 in mixtures of equal parts of Alcohol, Water and Ether, and in Alcohol, Glycerin and Ether. The formulas for various lotions and ointments are also given. — *P.J.* '05, 11 899

Petrosulfol—A dark brown thick syrupy substance. Soluble in Water. Similar in its therapeutic properties to Ichthyol.

Not Official

IGNATIA AMARA

The Seed of *Strychnos Ignatia*, Berg

Medicinal Properties—Similar in action to Nux Vomica.

Foreign Pharmacopœias—Official in Fr (Fève de St Ignace), Mex (Cabalonga), Port (Fava de S. Ignacio), Span (Haba de S. Ignacio)

EXTRACTUM IGNATIAE AMARAE—A dark brown liquid. Beans with Alcohol (90 p.c.), and evaporated to a thick extract. Tonic, given in debility of the digestive system.

Dosé — $\frac{1}{2}$ to 1 grain = 0.008 to 0.065 gramme in a pill three times a day.

Official in Mex

TINCTURA IGNATIÆ AMARÆ — 1 of Ignatia Beans, percolated with Alcohol (70 p c) to yield 10

Dose — 5 to 20 minims = 0.3 to 1.2 cc

Foreign Pharmacopœias — Official in Mex (Tintura de Caba-longas), 1 in 5

TEINTURE DE FÈVE DE SAINT-IGNACE COMPOSÉE (Fr). — St Ignatius Beans (rasped), 100, Potassium Carbonate, 2.5, Prepared Soot, 0.5, Alcohol (70 p c), 500, macerate for 10 days, and filter

INFUSA.

INFUSIONS

FR, APO/EMLS, TISANFS, GER, AUFGÜSSE, ITAL, INFUSI, SPAN, INFUSIONES

Infusions, though generally made with boiling Water, are in some cases ordered to be made at a lower temperature, as Infusum Calumbæ, the starch of which would be dissolved by boiling Water. The mucilage and vegetable albumen present are, however, dissolved by cold Water, and these render the Infusion liable to change.

When the Infusion is to be made with boiling Water the pot or vessel should be first rinsed with boiling Water. The ingredients should be suspended immediately under the surface of the Water, or otherwise should be stirred from time to time during infusion.

There is a very large demand for so-called Concentrated Infusions, but although very convenient and comparatively economical they have not the same characters as the freshly-made Infusions. B.P. '98 has included some *Liquores Concentrati* which are intended to represent Concentrated Infusions, they are fluid extracts, prepared with weak spirit (Alcohol 20 p c).

There are no General Directions given in the British Pharmacopœia for the preparation of Infusions.

General Directions given in German Pharmacopœia — For the preparation of Infusions boiling Water is poured on the medicament, which must be finely cut if necessary, heat for five minutes, with frequent shaking, on a water-bath, and strain after cooling. Infusions for which the amount of the respective substances is not specified, are prepared so that 10 parts of strained product are obtained from 1 part of substance. In the case of powerful substances for which a limit of dose is given, the quantity of substance is to be specified by the physician.

Directions in United States Pharmacopœia — An ordinary Infusion, the strength of which is not directed by the physician nor specified by the Pharmacopœia, shall be prepared as follows. Put 10 of the substance into a suitable vessel, provided with a cover, pour upon it 200 of boiling Water, cover the vessel lightly, and let it stand half an hour in a warm place, then strain and pass enough Water through the strainer to make the Infusion measure 200 parts. The strength of Infusions of energetic or powerful substances should be specially prescribed by the physician.

Two general methods are recommended by E. R. Farr and R. Wright for the preparation of Concentrated Infusions. They employ dilute Chloroform Water (1 in 1000) and Alcohol as a preservative, and the finished product when diluted in the proportion of 1 part to 7 parts of Water is fairly approximate to

the corresponding fresh Infusion. In the first process, **Repercolation**, half the drug is moistened with the menstruum and percolated, the remainder is then moistened and percolated with the first percolate until completely exhausted. The weak portions are evaporated and added to the stronger and made up to volume. By the second method that of **Macero-Expression**, the quantity of drug ordered per 20 fl oz is macerated in 15 oz of the menstruum in a covered earthenware vessel for 24 hours, slightly shaken when the drug is not completely covered with the menstruum, strain and press the marc, to the resulting liquid add any other ingredients specified, and reserve, repeat the maceration a second and third time for 6 hours each, and evaporate the resulting mixed liquors, add them to the reserved portion and make up to 20 fl oz, set aside for 7 days and filter. When diluted Alcohol is used the third maceration may be omitted, and only enough menstruum used in the second to make the expressed mixed liquids measure 20 fl oz. — *P. J. '06*, 163, 166, 169, 226, *U. S. '06*, 252, 253, *P. J. '07*, 1, 621, *Y. B. '07*, 247.

No. 33 Official

INULIN

ELI CAMPANI.

The Root of *Inula officinalis* L. It contains Inulin, a body altered to sugar, a crystalline bitter substance, Helenin or Alantcamphor. **Foreign Pharmacopœias** — Official in Mex. and Port. Not in the others.

HELENIN (C_8H_8O) — Colourless, acicular crystals, almost insoluble in Water, but readily soluble in hot Absolute Alcohol, Ether, and Volatile Oils. Has been found to possess powerful antiseptic properties, and has been given in bronchopneumonia, tuberculosis, and diphtheria.

Dose — $\frac{1}{2}$ to 2 grains = 0.016 to 0.13 gramme.

Official in Mex

IODIFORMUM.

IODOPHONUM TRI-iodomethane

Ca is a HI, eq 390 61

FR, IODOFORME, GER, IODOFORM, ITAL, IODOFORMIO, SPAN, YODOFORMO

The shining, lemon-yellow, low, small hexagonal crystals are official; but for dispensing purposes it is supplied as a fine crystalline **Powder**. There is also a **Precipitated Iodoform**, which, however, has a tendency to agglomerate. Iodoform has an unpleasant characteristic odour and taste, and is somewhat unctuous to the touch.

It should be kept in well-stoppered glass bottles of a dark amber tint in a cool atmosphere and should be protected as far as possible from the light.

It is chemically a tri-iodomethane and may be prepared by the action of Iodine and a reducing agent upon Ethyl Alcohol or alkali Carbonate upon Ethyl Alcohol.

Solubility — Very sparingly soluble in Water; 1 in 7 of Ether, 1 in 14 of Chloroform, 1 in 120 of Absolute Alcohol (90 per cent), and in debile volatile Oils.

1 in 30 of Olive Oil, 1 in 3½ of Carbon Bisulphide, sparingly in Petroleum Spirit

Precipitated Iodoform frequently gives a turbid solution in Chloroform and Carbon Bisulphide, owing to the dampness of the powder, the adhering Water being insoluble in those fluids. It rapidly dries on free exposure to air, and will then form a clear solution.

The above figures for solubility have been incorporated in the *B.P.C.* The note respecting the solubility of Precipitated Iodoform in Chloroform and Carbon Bisulphide appeared in the 15th edition of the *Compendium* and is aptly paraphrased in the *B.P.C.* as follows:—In the form of powder it sometimes contains a trace of moisture, consequently the solutions in Chloroform and Carbon Bisulphide may be turbid; a short exposure to the air, however, will quickly free it from the adherent moisture, when bright solutions may be obtained.

Medicinal Properties. Antiseptic, deodorant and local anæsthetic. Useful in cleansing foul ulcers, buboes, soft chancre, or syphilitic sores, the powder being applied, or an ointment (1 drm to 1 oz), or a solution in Oil of Eucalyptus. Used as a deodorant, and to relieve the pain of cancer and abate the progress of the disease, as a soothing application to burns, also to relieve neuralgia, goutic, and glandular enlargements, as a suppository in chronic proctitis, in hæmorrhoids and anal fissure.

A solution of Iodoform in Ether, containing about 10 p.c. of Liquid Paraffin in the proportion of ½ grain to 10 minims is used (*B.M.J.* '05, i, 67) as an intravenous injection in pulmonary tuberculosis. The Iodoform is placed in the syringe, the ether drawn in and shaken till the powder is dissolved. A few minims of Liquid Paraffin may be drawn up and the whole well shaken.

Intravenous injections are in pulmonary tuberculosis. *L.* '05, i, 1941.

A mixture of Iodoform, 60, Spermaceti, and Sesame Oil, of each, 40, has been used as a filling for chronic bone cavities. *B.M.J.* '05, i, 70.

An Ointment of Iodoform 5 to 10 grains; Vaseline, 2½ drms, or Hydrarg. Ox. Flav., 1 to 5 grains, Vaseline, 2½ drms, applied in keratitis. *M.P.* '05, ii, 308.

A case of presumably tuberculous meningitis successfully treated (*L.* '06, ii, 964) with an ointment containing 15 grains in an oz. of Vaseline thoroughly rubbed into the scalp and the back of the neck every 8 hours.

As an antiseptic, Iodoform in fine powder, alone or mixed with Boracic Acid or Bismuth, is used as an insufflation for ulcerated throat or for ozæna, and as a packing in bone cavities. *L.* '04, ii, 1317.

Whitehead's Varnish is Compound Finishing of Benzoin, in which Ether (sp. gr. 0.795) has been substituted for Alcohol (80 p.c.), and contains 10 p.c. of Iodoform.

To prevent pitting in smallpox (*L.* '86, ii, 89), injections of Iodoform in goitre (*L.* '81, 334), in tuberculous disease of the elbow joint. *B.M.J.* '97, ii, 897.

A stopping for bone cavities, Iodoform 30 to 60, Spermaceti, 40, Sesame Oil, 20. *B.M.J.* '01, ii, 16.

Daily hypodermic injections of 0.05 grammes of a mixture consisting of Iodoform, 15, Eucalyptol, 10, Liquid Vaseline 105, in recurrent hæmoptysis in the early stage of tuberculosis. *L.* '10, 705.

Dose. ½ to 3 grains = 0.032 to 0.196 grammes.

Ph. Ger. maximum single dose, 0.1 grammes, maximum daily dose, 0.6 grammes.

Prescribing Notes. The Iodoform should be finely powdered, or still better, precipitated Iodoform should be used, and suspended with Mucilage of Acacia for a mixture or lotion, or it may be given in pills made with Glucose or one sixth of its weight of Compound Powder of Tragacanth and Dispensing Syrup, or Diluted Glucose, q.s. to mass.

To cover the smell of Iodoform, Oil of Geranium (5 minims to 2 drms) answers

COLLODIUM IODOFORMI—Iodoform, 1, Flexible Collodion, 9—*Guy's*.

Official in Belg, Fr and Jap, 1 in 10 by weight

EMULSIO IODOFORMI—Iodoform, in fine powder, 10 parts, Glycerin, 70 parts, Water 20 parts Rub the Iodoform to a smooth paste with the Glycerin, then add the Water—*University*

This has been incorporated in the *B.P.C.*

Iodoform, 1, Glycerin, 10 p c, q s to moisten, Boiling Distilled Water, 2, Glycerin, 7—*Great Northern and Guy's*.

GLYCERINUM IODOFORMI—Iodoform (washed with 1 in 20 solution of Phenol to sterilise), 1, Glycerin, 9—*King's*

GOSSYPIUM IODOFORMI—Iodoform, 70 grains, Glycerin, 10 minims; Cotton-Wool 60 grains, Ether and Absolute Alcohol are used as solvents. Contains about 50 p c of Iodoform

Iodoform Wool can also be obtained containing 10 p c and 20 p c Iodoform Lint 10 p c

Official in Austr 10, 20, and 30 p c, Belg, Dutch and Span. 10 p c; Jap 5 p c

INJECTIO IODOFORMI—Iodoform, 1, Mucilage of Tragacanth, 2, Water, 7—*University*

This has been incorporated in the *B.P.C.*

Saturated Solution of Iodoform in Ether, $\frac{1}{2}$ fl oz, Olive Oil, to 1 fl oz—*Central Throat*

INSUFFLATIO IODOFORMI—(For throat) Iodoform, 2, Dried Starch, 1, both in fine powder (Aural) Iodoform, 1, Boric Acid, 3, both in fine powder.—*Throat*

Iodoform, in fine powder, 1, Subnitrate of Bismuth, 1.—*Throat* (1894).

This has been incorporated in the *B.P.C.*

INSUFFLATIO IODOFORMI ET MORPHINÆ COMPOSITA—Iodoform 1 grain, Boric Acid, 1 grain, Morphine Acetate, $\frac{1}{2}$ grain, Starch, to make 5 grains—*Guy's*

NEBULA IODOFORMI—Iodoform, 40 grains, Ether (sp gr 0.735), 1 fl oz; dissolve. A strong antiseptic and detergent

Iodoform, 8, Ether, q s to produce 100—*B.P.C.*

LIQUID IODOFORM—Caustic Potash, 35, Water, 25, dissolve, shake well, and add Oleic Acid 50, Alcohol (95 p c), 30, then add with continuous agitation Sublimed Iodine, 30, decolorise by the addition of a few drops of solution of Caustic Potash. Set aside for several days in the dark, and decant the supernatant liquid. A yellowish liquid, with the odour of Iodoform, miscible with Water, Alcohol, Ether, and Chloroform, is thus obtained (*Blanchi*)—*P.J.* 07, 1 509, '06, 1 663, *C.D.* '07, 1 382; '06, 1 163

PASTILLUS IODOFORMI—Iodoform, in fine powder, 1 grain, Glycerin, 1 minim; Glyco-gelatin, 18 grains For one pastille

PIGMENTUM IODOFORMI—Iodoform, 1, Ether, 8—*Central Throat*.

PULVIS IODOFORMI COMPOSITUS. *S. Nav.* Iodoform.—Iodoform, 20, Boric Acid, 30, Naphthalene, 50, cf *Fr.* 11 25 This powder is used in many cases where a diluted preparation of Iodoform, for external purposes, is desired—*U.S.N.F.*

Iodoform, in fine powder, 1, Boric Acid, 3. For external use only.—*East London for Children and St. Thomas's*

This has been incorporated in the *B.P.C.*

UNGUENTUM IODOFORMI CUM ATROPINA—Precipitated Iodoform, 60 grains, Atropine, 2 grains, Soft Paraffin, 1 oz, heat the Atropine and Paraffin till dissolved, stir, and while cooling add the Iodoform—*London Ophthalmic and St. Mary's*

This has been incorporated in the *B.P.C.*

UNGUENTUM IODO-PARAFFINI.—Iodoform, 1, Eucalyptus Oil, 8, Soft Paraffin, 27, Hard Paraffin, 6

Dissolve the Iodoform in the Oil at a slightly raised temperature, and mix with the other ingredients previously melted together — *University* (1899)

Unguentum Iodoformi et Eucalypti — Iodoform, 2, Oil of Eucalyptus, by weight, 19, Hard Paraffin, 64.50, Soft Paraffin, 14.50 — *B P C*

VASOLIMENTUM IODOFORMI — Iodoform, 1.5, Liquid Vasoliment, 98.5 — *Hager*

Parogenum Iodoformi *Syn* Iodoform Vasoliment — Iodoform, 3, Parogen, *q s* to produce 100 — *B P C*

Iodoform, 1.5, Parogen, 98.5 — *P J '06*, i 619, *Y B P '06*, 147

VASOLIMENTUM IODOFORMI DESODORATUM. — Iodoform, 1.5, Eucalyptol, 1.5, Liquid Vasoliment, 97 — *Hager*

Parogenum Iodoformi Desodoratum *Syn* Deodorised Iodoform Vasoliment — Iodoform, 3, Eucalyptol, 3, Parogen, *q s* to produce 100 — *B P C*

Iodoform, 1.5, Eucalyptol, 1.5, Parogen, 97 — *P J '06*, i 619, *Y B P '06*, 147

Dissolve the Iodoform in the Parogen by warming cautiously, and add the Eucalyptol

EKA-ODOFORM — A yellow-lemon crystalline lustrous powder, insoluble in Water. Soluble 1 in 75 of Alcohol (90 p c), 1 in 8 of Ether, 1 in 18 of Chloroform. Stated to be a mixture of Iodoform and Paraformaldehyde. Introduced as a substitute for Iodoform

Iodofan — A reddish crystalline powder without taste or smell, possessing anti-bacterial and deodorising properties, introduced as a dressing — *B M J E '07*, ii 63

ODOFORMIN — A combination of Iodoform and Hexamethylenetetramine containing about 75 p c of the former. A white or pale yellow powder, insoluble in Water, soluble in 1 in 170 of Alcohol (90 p c), 1 in 850 of Ether, 1 in 72 of Chloroform, also soluble in Acetone. Boiling Water, Acids, and Alkalis decompose it. Introduced as an Iodoform substitute — *J S C I '95*, 820, '96, 469, '97, 757, *C D '95*, ii 488; *P J '95*, ii 455, '97, ii 82, *L '96*, i 856

Iodoformal (Iodoformin Ethyl Iodide) — In yellow crystals or powder, insoluble in Water. Antiseptic

ODOFORMOGEN (Iodoform Albuminate) — A pale lemon yellow powder, possessing a faint odour of Iodoform. Insoluble in Water, Alcohol (90 p c), Ether or Chloroform. Used as a dusting powder. Introduced as an Iodoform substitute — *B M J '98*, ii 1066 *B M J E '98*, ii 63

DI-ODOFORM (Ethylene Periodide) — Yellow prismatic needles. Insoluble in Water, soluble in Chloroform. Introduced as a substitute for Iodoform — *L '98*, ii 1355, *Pr* li 126, *P J*, (3) xxiv 622. It is official in *F. Codex* (1908)

IODOL Tetranodpyrrol C₄H₄NH₄, eq 566.18 — A light brown microcrystalline powder, without taste, having a faint odour, and containing 90 p c of Iodine. It should be kept in well closed glass bottles of a dark amber tint in a cool place and protected as far as possible from the light

Solubility — Nearly insoluble in Water, 1 in 18 of Alcohol (90 p c), 1 in 150 of Chloroform, 1 in 1½ of Ether, 1 in 155 of Glycerin. It is stated to be soluble 1 in 3 in Absolute Alcohol, but the sample we examined gave 1 in 6½

Medicinal Properties — Antiseptic, used for the same purposes as Iodoform, but it is free from the objectionable odour of the latter, and is stated not to be so poisonous. 1 p c of Menthol is added in nasal dressings to cover the odour of Iodol

Foreign Pharmacopœias. — Official in Ital., Mex., Russ., Span., and U.S.; not in the others

Formulae. — *Yodol* (Mex.), Iodol, 1, Vaseline, 9

Tests—Iodol does not undergo decomposition when heated up to 100° C (212° F), but at about 130° C (302° F) it is decomposed, giving violet coloured vapours of Iodine. When warmed with 10 p.c. Sodium Hydroxide Solution and Zinc foil, it evolves vapours of Pyrrol, which impart a bright red to a deep carmine-red colour to a splinter of pine wood moistened with Hydrochloric Acid.

A weighed quantity of 0.5 gramme should leave no weighable residue when ignited with free access of air. The *U.S.P.* states that, when ignited, it should leave not more than 0.1 p.c. of residue, the limit of inorganic impurities or mineral residue. When shaken with Water and filtered the filtrate should yield not more than the slightest opalescence with Silver Nitrate Solution, indicating the limit of Hydrochloric Acid and soluble Iodides, nor any coloration with Hydrogen Sulphide, indicating the absence of heavy metals, *e.g.*, Copper, Lead. When Water which has been shaken with the sample is in turn shaken with Carbon Bisulphide the latter should be coloured at the most a pale yellow, not a violet.

IODOLENE (Iodol) light yellow powder, insoluble in Water and in Alcohol (90 p.c.) forms, one for internal use containing 10 p.c. Iodol, the other for external use containing 36 p.c. Antiseptic. Introduced as an Iodoform substitute. Has been used internally in syphilis, but has sometimes caused iodism. — *B.M.J.* '02, 1 91

Dose—15 to 30 grains = 1 to 2 grammes.

IODUM.

IODINE

I, eq 125 90

FR, IODE SUBLIMÉ, CLR, JOD, ITAL, JODO, SPAN, YODO

Heavy, greyish-black, rhombic plates or prisms, possessing a metallic lustre and a characteristic peculiar odour. Commercial resublimed Iodine, if in large dry scales, may be reckoned at 100 p.c. It is prepared from 'kelp' (the ashes of sea-weeds), and also from naturally occurring Iodides and Iodates. It should be kept in well-stoppered glass bottles and in a cool atmosphere, as it volatilises considerably at ordinary temperatures.

Solubility—1 in 7000 of Water, 1 in 12 of Alcohol (90 p.c.); 1 in 4 of Ether, 1 in 30 of Chloroform, 1 in 6 of Carbon Bisulphide, 1 in 65 of Glycerin, soluble in an aqueous solution of Potassium Iodide.

Medicinal Properties—Antiseptic, alterative, deodoriser, disinfectant, locally it is irritant or vesicant according to the strength employed. Internally, largely used in form of Iodide, seldom as Iodine, in chronic rheumatism and in chronic inflammation of various kinds, to promote absorption in hepatic and splenic enlargements, and in dropsies (pericardic effusion, hydrocele, etc.) In the form of Potassium Iodide (10 to 30 grains three times a day), it is specific in the later stages of syphilis, and in 30-grain doses three times a day it is very useful in aneurism, its most striking effect being the relief of the aneurismal pain; valuable in actinomycosis. Efficacious in all chronic inflammatory conditions, caution, however, is required, as it may, when given in very large doses, occasionally cause wasting

of healthy glands, such as the mammæ and testes 1 of the Tincture with 50 of Water forms an antiseptic lotion for washing out cysts Externally the solution, ointment, and tincture are applied in chronic and parasitic skin diseases, in phthisis, pleurisy, pericarditis and bronchitis as a counter-irritant, and for chilblains; the Tincture, either neat or diluted with an equal quantity of Water, is injected into the scrotal sac to cure hydrocele, Morton's Fluid is injected into the sac of spina bifida A few drops of the Tincture in half a pint of hot Water may, along with Creosote or volatile Oils, be inhaled in some forms of chronic bronchitis and phthisis, and in the throat affection of scarlatina and measles It is employed as a gargle, 1 or 2 of Tincture in 32 of Water for ulceration of the throat. One or two drops of the Tincture in a tablespoonful of Water every 30 minutes are often successful in checking vomiting, including that of pregnancy See also under 'Potassii Iodidum'

Half a syringe of a solution of Iodine 1, Potassium Iodide 2, and Water 50, injected into the genital region in tuberculous peritonitis — *B M J F* '99, ii, 44.

Stated (*P J*, '04, ii 987) to form an excellent general tonic before meals in tuberculosis, a teaspoonful of the following mixture being recommended Tincture of Iodine (*Fr Codex*), 20, Potassium Iodide, 2, Glycerin, 40, Syrup of Orange, 50, Water, to 1000 1 minim dose of the Tincture (*B M J* '04, ii, 1405), very successful in sea sickness

For the relief of troublesome cough of phthisis Dr. Coghill's famous formula (*Edin Med Jour* '05, i 465) is useful — Tincture of Iodine (Ethereal), 2 drm, Acid Carbohc, 2 drm, Creosote or Thymol, 1 drm, Alcohol (90 p c) to 1 oz

Equal parts of the liniment and tincture applied as a paint in pleurisy of phthisis — *Edin Med Jour* '05, i 468

A case of acute poisoning with fatal result caused by drinking 4 oz of Liniment of Iodine — *L* '05, i 798

In form of tincture, strongly recommended in carbolic acid poisoning, in large doses, possibly up to drm doses and more in severe cases — *L* '07, ii 298

Graves' disease treated with marked success by parenchymatous injections of Iodine and Ergotine — *B M J F* '06, ii 87 If employed from the beginning in typhoid, it acts almost like a specific, shortening the duration of the illness, and modifying favourably most of the symptoms Given as *B P* tincture 8 to 16 minims in 1 or 2 fl drm of Rum or Cognac in 1 or 2 oz of Water with Sugar, 8 or 4 times in 24 hours — *B M J* '07, ii 148

Dose.— $\frac{1}{4}$ to $\frac{1}{2}$ gram = 0.001 to 0.015 gramme

Ph *Gr* maximum single dose, 0.02 gramme, maximum daily dose, 0.06 gramme

Prescribing Notes — Iodine rarely given internally in the solid form, except when loosely combined as in the alkaloidal Periodides, see p 276 Occasionally administered as Tincture, which should be well diluted The *Pasta Iodii et Amyli* is less irritating than many of the other Iodine preparations

Iodine and solutions containing free Iodine stain the skin a yellowish brown; this can be removed by Caustic or Carbonated Alkali or Sodium Thiosulphate Several so-called colourless and non-staining preparations of Iodine have been suggested, but their medicinal action cannot be due to free Iodine, but to the compound of Iodine which is produced in each case, e.g., combinations of Iodine and Oleic Acid and the fixed and volatile Oils, also the Decolorised Tincture of Iodine (*B P C*) which is practically a solution of Iodine and Potassium Iodide

In all the galenic preparations containing Iodine, Potassium Iodide is a constant ingredient, presumably with the intention of assisting the solution of the Iodine In the case of aqueous solutions this is necessary, and an excess of Iodide is advantageous In spirituous solutions, however, where the Iodide is scarcely more soluble than the Iodine, a much smaller quantity (if any) is required

decolorise the free Iodine, the *PG* Sulphurous Acid, the *USP* uses Ferrous Sulphate Solution and Sodium Hydroxide Solution (5 p.c.), the *PG* a crystal of Ferrous Sulphate, a drop of Ferric Chloride Solution, and Sodium Hydroxide Solution (15 p.c.) both Pharmacopœias require that no blue coloration should be produced on the addition of a slight excess of Hydrochloric Acid, Chloride and Bromide would appear in the aqueous extract, and are precipitated with Silver Nitrate Solution, the precipitated Iodide remains insoluble when treated with Ammonia Solution, the Chloride and portion of the Bromide if present passing into the ammoniacal solution, the latter is required to yield not more than a slight opalescence, and certainly no precipitate when rendered slightly acid with Nitric Acid

Ferrous Sulphate and Sodium Hydroxide—Triturate 0.5 gramme of finely powdered Iodine with 20 c.c. of Water and filter the solution. To one half of this solution in a test tube carefully add Tenth normal Volumetric Sodium Thiosulphate Solution, until the solution is just decolorised. Then add a few drops of Ferrous Sulphate T.S. and subsequently a little Sodium Hydroxide T.S. and heat the mixture gently. On now adding a slight excess of Hydrochloric Acid the liquid should not assume a blue colour (absence of Iodine Cyanide), *USP*, triturate $\frac{1}{2}$ gramme of the sample with 20 c.c. of Water, filter and decolorise a portion of the filtrate with Sulphurous Acid. Warm a portion of the decolorised liquid with a crystal of Ferrous Sulphate, a drop of Ferric Chloride T.S. and Sodium Hydroxide Solution (15 p.c.) This mixture shall yield no blue coloration on the addition of an excess of Hydrochloric Acid, *PG*

Silver Nitrate and Ammonia Solution—To the other half of the aqueous filtered solution obtained in the preceding *USP* test add a slight excess of Silver Nitrate T.S., shake the liquid actively, allow the precipitate to subside, and having poured off the supernatant liquid completely, shake the precipitate with a mixture of 1 c.c. of Ammonia Water and 9 c.c. of Water, and filter. Upon the addition of a slight excess of Nitric Acid to the filtrate not more than a slight opalescence should make its appearance, *USP*, add to the portion of the filtrate remaining from the above *PG* test an excess of Ammonia Solution and of Silver Nitrate Solution, and filter. The filtrate when acidified with Nitric Acid shall yield at most an opalescence, but not a precipitate, *PG*

Volumetric Determination—A solution of 1 gramme of Iodine and 2 grammes of Potassium Iodide in 50 c.c. of Water should require for decolorisation at least 78.4 c.c. of Volumetric Solution of Sodium Thiosulphate, *BP*, 0.2 gramme of Iodine and 1 gramme of Potassium Iodide dissolved in 20 c.c. of Water should require for decolorisation at least 15.6 c.c. of Tenth normal Volumetric Solution of Sodium Thiosulphate, *PG*. About 0.5 gramme of Iodine is accurately weighed and dissolved with 1 gramme of Potassium Iodide in 50 c.c. of Water and titrated with Tenth normal Volumetric Sodium Thiosulphate Solution until the solution is decolorised. The number of c.c. of the Volumetric Solution required when multiplied by 1.269 and divided by the weight of Iodine taken gives the percentage of pure Iodine present, *USP*

Preparations.

LIQUOR IODI FORTIS. STRONG SOLUTION OF IODINE
LINIMENT OF IODINE, *BP*. '85

Iodine, $1\frac{1}{2}$, Potassium Iodide, $\frac{3}{4}$; Distilled Water, $11\frac{1}{2}$, Alcohol (90 p.c.), 9. (about 1 of Iodine in 8 $\frac{1}{2}$)

Formerly called Linimentum Iodi Alcohol (90 p.c.) and Distilled Water replace the Rectified Spirit of Glycerin The Potassium Iodide is increased

Foreign Pharmacopœias—Official in Dutch (Solutio Lugoli), Iodine 1, Potassium Iodide 2, Water 497, Norw (Solutio Supræiodetæ Kalici), Iodine 1, Potassium Iodide 2, Distilled Water 97, Port (Solutio Iodo-iodetado), Tincture of Iodine 6, Potassium Iodide 1, Water 18, U.S. (Liquor Iodi Co), Iodine 1, Potassium Iodide 2, Distilled Water 17 All by weight Not in the others

Tests.—Strong Solution of Iodine has a sp gr of 1.008 to 1.012; contains 11.7 p.c. w/v of Iodine as determined by titration with Volumetric Sodium Thiosulphate Solution, about 6.8 p.c. w/v of total solids and 65 p.c. w/v of Absolute Alcohol as determined by the distillation method given under Tinctura Iodi

TINCTURA IODI. TINCTURE OF IODINE

Iodine, $\frac{1}{2}$, Potassium Iodide, $\frac{1}{2}$, Distilled Water, $\frac{1}{2}$, Alcohol (90 p.c.), q.s. to yield 20 (1 of Iodine in 40)

The Iodine and Iodide are first dissolved in a small quantity of Water, as suggested in previous editions of the *Companion*

Dose.—2 to 5 minims = 0.12 cc to 0.3 cc

Ph. Ger. maximum single dose, 0.2 gramme, maximum daily dose, 0.6 gramme of the 1:10 tincture

Tinctura Iodinei (*Ph. Ger.*) 1 of Iodine in 16 of Alcohol (90 p.c.) It resembles the Tinctures Pharmacopœias in being without Potassium Iodide

Tinctura Iodi Ætherea (*Swayer*)—1 of Iodine in 40 of pure Ether.

Foreign Pharmacopœias—Official in Austr, Belg, Dan (Solutio Iodi Spirituosa Concentrata), Dutch (Solutio Iodii Spirituosa), Fr, Port and Swiss, 1 and 9 Ital, Jap and Mex, 1 and 12, Mex has (Tintura de Yodo Yodurado), Potassium Iodide 1, Tincture of Iodine 1, Norw and Swed (Sol Iodii Spirituosa) 1 in 20, Swed also includes Solutio Iodii Concentrata 1 in 10 Ger, Hung, Russ and Span 1 and 10, U.S., Iodine 7, Potassium Iodide 5, Alcohol to 100

Tests—T. c. . . . has a sp gr of 0.875 to 0.880. It is officially required to contain 2.47 p.c. w/v of Iodine as determined by titration with Volumetric Sodium Thiosulphate Solution. It contains about 2.5 p.c. w/v of total solids and about 86.0 p.c. w/v of Absolute Alcohol. Before determining the Alcohol by distillation it is necessary to fix free Iodine. This may be accomplished by the addition of Sodium Hypo-sulphite Solution (50 p.c.), the liquid being then neutralised with Potassium or Sodium Hydroxide

UNGUENTUM IODI. IODINE OINTMENT

Iodine, 20 grains, Potassium Iodide, 20 grains, Glycerin, 60 grains, Lard, 400 grains (1 of Iodine in 25)

B.P. 1835 was 1 in 31

Foreign Pharmacopœias—Dutch, Iodine 2, Potassium Iodide 3, Water 5, Ointment 90, Fr (Pommade d'Iodure de Potassium Ioduré), Iodine 1, Potassium Iodide 5, Benzoated Lard 40, Water 4, Hung, Tincture of Iodine 1, Simple Ointment 10, Mex (Pomada de Yodo), Iodine 1, Lard 80; Port (Pomada de Iodeto de Potassio Iodada), Iodine 1, Potassium Iodide 4, Water 5, Lard 40, Span (Pomada de Ioduro Potassico Iodado), Iodine 2, Potassium Iodide 2, Glycerin 6, Lard 40, U.S., Iodine 4, Potassium Iodide 4, Glycerin 12, Benzoinated Lard 80 Mix. Not in the others.

Not Official

CAUSTICUM IODI —Iodine, 180 grains, Potassium Iodide, 60 grains, Alcohol (90 p c), 1 fl oz

Used in cases of lupus and of indolent (i e non phagedenic) tertiary syphilitic ulcers

CHLOROFORMUM IODI —Iodine, 1, Chloroform, q s to produce 10 —
Martindale

This has been incorporated in the *B P C*

COLLODIUM IODATUM (U S N F) —Iodine, 1, Flexible Collodion, 19

Collodium Iodi —Iodine, 6 50, Acetone Collodion, q s to produce 100 —
B P C

GOSSYPIUM IODATUM —Dry white wool impregnated with Iodine, and containing about 8 p c of the latter (Cotton Iodid, *Fr Codex*, at least 4 p c).

INHALATIO IODI C CONIO — $\frac{1}{2}$ to 1 fl drm of Succus Conii added to Vapor Iodi

GLYCERINUM IODI (Morton's Fluid) —Iodine, 10 grains, Potassium Iodide, 30 grains, Glycerin, 1 fl oz —*Guy's*

For spina bifida, inject 30 minims, without allowing the fluid contents of the tumour to escape —*B M J* '85, 1 1098, '86, '874, '87, 11 1275

Liquor Iodi Glycerinus (Morton's) —Iodine, 10 grains, Potassium Iodide, 30 grains, Glycerin, 1 oz Dissolve —*Pharm Form*

Note —It is advisable to dissolve the Iodine and Iodide in about $\frac{1}{2}$ drm of Water before adding the $\frac{7}{8}$ drm of Glycerin —*Pharm Form*

This has been incorporated in the *B P C*, as follows —**Glycerinum Iodi**
Syn **Injectio Iodi**, Morton's Fluid —Iodine, 2, Potassium Iodide, 6, Distilled Water, 5, Glycerin, q s to produce 100 —*B P C*

INJECTIO IODI —Solution of Iodine, 1 fl drm, Water, to 20 fl oz —
Samaritan

PHENOL IODATUM See p 36

LUGOL'S CAUSTIC —Iodine, 1, Potassium Iodide, 1, Water, 2.

LUGOL'S SOLUTION —Iodine, 20 grains; Potassium Iodide, 30 grains, Water, 1 oz This was official as **Liquor Iodi** in *B P* '85, but omitted in '98 The proportions are about equal to 1, $\frac{1}{2}$, and 22 ^o See also **Liquor Iodi Fortis**

LIQUOR IODI (*B P* '85) —Iodine, 10, Iodide of Potassium, 15, Distilled Water, q s to produce 200

This has been incorporated in the *B P C* under the title **Liquor Iodi Dilutus**.

LIQUOR IODI COMPOSITUS (U S P) —Iodine, 5, Potassium Iodide, 10; Distilled Water, q s to make 100 by weight

LIQUOR IODI CARBOLATUS *Syn* **Bonillon's Solution**, French Mixture —Compound Solution of Iodine (*U S P*), 15, Carbohc Acid, liquefied by gentle heat, 5 5, Glycerin, 165, Water, q s to make 1000 *U S N F*

NEBULA IODI COMPOSITA —Iodine, 1 grain, Carbohc Acid, 4 grains, Spray Oil, 1 fl oz —*Bournemouth Formulary*

Iodine, 1, Carbohc Acid, 1, Liquid Paraffin, q s to produce 100 —*B P C*

NEBULA IODI ET MENTHOLIS —Iodine, 1 grain, Menthol, 1 drm, White Petroleum Oil, q s to make 1 fl oz —*A Ph F*

Iodine, 2, Menthol, 4, Liquid Paraffin, q s to produce 100 —*B P C*

PASTA IODI ET AMYLI —Starch, 1 oz, Glycerin, 2 fl oz, Water, 6 fl oz, boil together, and when nearly cold add Solution of Iodine, *B P* '85, 1 fl oz. —*University*

This has been incorporated in the *B P C*

PIGMENTUM IODI —Iodine, 2, Potassium Iodide, 1, Glycerin, 4. Used to destroy vegetable parasites

Tincture of Iodine, 1, Strong Solution of Iodine, 1 — *Great Northern, Middlesex, University*

This is equivalent to 1 in 24 of Iodine; most of the Hospitals have a Pigmentum, varying in strength from 1 in 8 to 1 in 84, some with Glycerin, others without. Pigmentum Mandl is 1 in 73.

Iodine, 100, Potassium Iodide, 100; Water, to 1 fl oz — *St Thomas's*.

This has been incorporated in the *B P C*

PIGMENTUM IODI CUM ACONITO MITE — Tincture of Iodine, 1; Tincture of Aconite, 1 — *R D H*

PIGMENTUM IODI CUM ACONITO FORTE — Strong Solution of Iodine, 1, Liniment of Aconite, 1 — *R D H*

PIGMENTUM IODI CARBOLISATUM — 4 grains, Iodide of Potassium, 4 grains, Carbolic Acid, 4 4 fl drms, Water, to 1 fl oz

Dissolve the Iodine and the Iodide of Potassium in the Water, and then add the Carbolic Acid dissolved in the Glycerin

Note — This is sometimes employed at half strength — *Central Throat*.

This has been incorporated in the *B P C*

Pigmentum Iodi Oleatum — Iodine, 50 grains, Oleic Acid, to 1 fl oz — *Central Throat*

PIGMENTUM MANDL — Iodine, 6 grains, Potassium Iodide, 20 grains; Oil of Peppermint, 5 minims, Glycerin, to 1 fl oz — *Throat* Use, in granular pharyngitis

In answer to an inquiry for the correct composition of Mandl's solution, several formulas were given. That given in *Hager* is Carbolic Acid and Iodine, of each, 1, Potassium Iodide, 2, Glycerin, 100, but all the formulas sent in reply omitted the Carbolic Acid — *P J C*, 156, 181, 184, 200

PIGMENTUM PICIS CUM IODO (Coster's Paste) — Iodine, 120 grains; Rectified Oil of Tar, 1 fl oz — *Middlesex* dissolve cautiously, applying a gentle heat as required. The Oil of Tar is inflammable.

Specially recommended in ringworm.

This has been incorporated in the *B P C*

SIROP IODOTANNIQUE (Fr) — Iodine, 2, Tannin, 4, Distilled Water, 860, Refined Sugar, 640, all by weight.

Powder the Iodine and Tannin, with the Tannin and Water, into a flask, which heat on a water-bath at a temperature of 60° C until a drop of the solution ceases to give a blue colour on blue paper, then dissolve the Sugar in the solution

SIROP IODOTANNIQUE PHOSPHATÉ (Fr) — Monocalcic Phosphate, 2, Iodotannic Syrup, 98

TINCT IODI (P L) — Iodine 2½ Rectified Spirit 40 Dissolve the Iodine in the Spirit with the aid of a gentle heat and agitate

TINCTURA IODI DECOLORATA — Iodine, 250 grains, 250 p c, 1, 5½ fl oz, dissolve with a gentle heat, when cold add Strong Solution of Ammonia, 10 fl drms, keep the mixture in a warm place until decolourised, after which dilute with Alcohol (940 p c) to make 20 fl oz — *B P C Formulary 1901* incorporated in *B P C* as follows — Iodine, 2 30, Strong Solution of Ammonia, 6 25, Alcohol, 9 5 to produce 100

Liquor Ammoniae Iodidi (Simpson) — Liq Ammon Fortis, 2 fl oz; Iodine, 10 grains, Potassium Iodide, 20 grains, Alcohol (90 p c), 1 fl. oz; dissolve

Tinctura Iodi Decolorata — Iodine, 83, Sodium Thiosulphate (*U S P*), 88, Water, 100, Stronger Ammonia Water (*U S P*), 65, Alcohol (95 p c), 9 5 to produce 1000 — *U S N F*

UNGUENTUM IODI DENIGRESCENS *Syn.* Stainless Iodine Ointment — Iodine, 1 oz Soft Paraffin, 19 oz Powder the Iodine, melt the Paraffin by heat, add the Iodine and continue to heat the mixture, stirring until the Iodine

is combined Remove the heat and stir the preparation until cold—*Canadian Formulary*, also in *Pharm Form*

This has been incorporated in the *B P C*

VASOLIMENTUM IODI—Iodine, 10.5, Oleic Acid, 50, Alcoholic Ammonia, 25, Liquid Paraffin, 100, after solution is effected The weight is then made up to 175 with Alcohol—*Pharm Codex* *ALI* 756, *B P* '01, 212

Parogenum Iodi *Syn* Iodine Vasoliment—Iodine, 10, Oleic Acid, 40, Liquid Paraffin, 40, Ammoniated Alcohol (10 p c), 10 *B P C*

VASOLIMENTUM IODATUM—Iodine, 6, Liquid Vasoliment, 94—*Hager*

Parogenum Iodi Dilutum *Syn* Diluted Iodine Vasoliment—Iodine Parogen, 6, Parogen, 4—*B P C*

VAPOR IODI (Inhalation of Iodine)—Tincture of Iodine, 1 fl drm; Water, 1 fl oz mix in a suitable apparatus, and having applied a gentle heat, let the vapour that arises be inhaled This was official in *B P* '67 and '85, but omitted in '98, and has now been incorporated in the *B P C*

Tincture of Iodine 10 drops for each dry inhalation, without the aid of heat

COGHILL'S INHALATION FLUID—Iodine, 83 grains, Ether, 8 fl drm, Carbolic Acid, 8 fl drm, Creosote (or Thymol), 4 fl drm, Rectified Spirit, to 4 fl oz—*Pharm Form* (1 in 58)

VAPOR IODI ÆTHEREALIS—Iodine, 3 grains, Ether, 2 drm, Carbolic Acid, 2 drm, Creosote, 1 drm, Alcohol (90 p c), 3 drm Thymol may be substituted for Creosote—*Martindale* (1 in 146)

Vapor Iodi Ætherealis—Iodine, 0.05, 1 Ether, 25, Carbolic Acid, 25, Creosote, 12.50, Alcohol, 37.50—*B P C* (1 in 2000)

Altered in *B P C Supplement* from 0.05 of Iodine to 0.086, or prepared by mixing Ethereal Tincture of Iodine, 25, Carbolic Acid, 25, Creosote, 12½, and Alcohol (90 p c), q s to make 100

VAPOR IODI ET ACIDI CARBOLICI—Tincture of Iodine, 2 fl drm, Carbolic Acid, 2 drm, Thymol, 1 drm, Chloroform, 30 minims, Alcohol (90 p c), to produce 8 fl drm 10 to 20 drops twice or three times daily on a dry inhaler—*King's*

VAPOR IODI COMPOSITUS—Tincture of Iodine, ½ oz, Creosote, 1 fl drm, Liquefied Phenol, 1 fl drm, Rectified Spirit, ½ fl oz For dry inhalation—*Great Northern*

IODINE LEAF—An ingenious method for the local application of Iodine as a counter irritant, being two sheets of filter paper, one saturated with a solution of Potassium Iodide and Iodate, and the other with Acid Potassium Sulphate When the papers are moistened and brought together, Iodine is liberated—*L* '02, 1 328

Sajodin—A white powder, free from smell or taste, stated to contain 26 p c of Iodine, Dose, 2 to 3 grammes daily in syphilis—*M J E* '07, 1, 64

Iodalbin—Iodine in combination with Albumen, stated to contain 21.5 p c of Iodine, a reddish powder, insoluble in Water and Alcohol, but is dissolved by alkaline solutions Dose, 5 grains

IODI TRICHLORIDUM Iodine Trichloride or ICl_3 , eq 231.47—Orange yellow crystalline masses evolving a powerful penetrating chlorinous odour It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light

Solubility—1 in 1 of Water, 1 in 1 of Alcohol (50 p c)

Powerful antiseptic and disinfectant

Tests.—Iodine Trichloride melts at about 25° C (77° F) When heated with Oxalic Acid it yields violet coloured vapours of Iodine

The 1 in 10 aqueous solution affords on the addition of a considerable excess of Sulphuric Acid a whitish precipitate, changing to yellow

It contains theoretically 54.89 p c of Iodine A weighed quantity of 0.1 gramme mixed with 2 grammes of Potassium Iodide, and dissolved in 80 c.c. of Water, requires at least 15.0 c.c. of Decinormal Volumetric Sodium

Thio-sulphate Solution for decolorisation. A weighed quantity of 0.1 gramme should leave no weighable residue when ignited with free access of air.

IODIPIN—Under this title two Iodine addition-compounds of Sesame Oil are known: one containing 10 p.c. Iodine, a pale straw coloured transparent oily fluid, the other 25 p.c. a yellowish-brown viscid fluid. Both possess an oleaginous odour and taste, are insoluble in Water and in Alcohol (90 p.c.), soluble in all proportions of Ether and of Chloroform.

Medicinal Properties—Recommended in conjunctivitis, bronchial asthma, emphysema and pleuritis, also in ophthalmia of the eye—*B.M.J.E.* '00, ii 80, '01, ii 79, *M.A.* '02, 38.

As a test for the functional activity of the stomach—*B.M.J.E.* '99, ii 81.

In the treatment of fibroids growing at the menopause, Iodine by the mouth used to be successfully employed, the modern idea is in favour of the employment of hypodermic injections of Iodine in organic combination. Iodipin has been shown (*B.M.J.* '04, ii 1085) to give rise to some cases. The question of the originality of this hypodermic method has given rise to a good deal of correspondence (*M.P.* '04, ii 177, '05, i 11). In general, however, the results of the treatment have been injected every other day in the region of the affected parts.

No case of tertiary ulceration should be despaired of until Iodipin has been tried, its full effect secured by oral administration, is well borne, produces no ill effects.—*E.T.* '07, 45.

Prescribing Notes—May be given in capsules, each containing 2 grammes = 80 grains of the 25 p.c. preparation, or with Gum Acacia and flavoured with Oil of Cinnamon or Peppermint. It may also be administered subcutaneously, as in the treatment of syphilis, or byunction.

Dose—1 to 6 fl. drms = 3 G to 21 O cc of the 10 p.c. solution administered by the mouth. 150 to 300 grains = 10 to 20 grammes administered daily subcutaneously.

In cases of uterine fibroid, starting with 1 cc injected into the buttock and increasing up to 5 and 7 cc of the 25 p.c. injected into the buttock alternately, then 10 cc of the 10 p.c. and finally 10 cc of the 25 p.c., which latter is considered the full dose of Iodipin for ten consecutive days' use.—*L.* '03, i 959.

IOTHION $C_{12}H_{10}I_2O_2$, eq 309.41—This substance forms a pale yellow syrupy liquid, insoluble in Water, soluble in Alcohol (90 p.c.), Chloroform and Ether, and mixes with most oils and fats. It contains about 80 p.c. of Iodine, and being absorbed with facility by the skin, it forms a ready means of administering Iodine. Occasionally it is used in the treatment of the joints. It has been used successfully in poisoning by arsenic, and in the treatment of the joints. It may be painted on the skin, either pure or diluted with an equal quantity of oil, or may be used in the form of an ointment (25 to 50 p.c.) 80 to 60 grains per day is regarded as the average dose of pure Iothion.

For the gradual application of Iodine, it appears (*E.T.* '05, ii 28) to be of great value. It may be applied in the form of a 25 to 50 p.c. ointment made with Lanolin and Vasoline.

Prescribers should have their attention drawn to the fact that it possesses a sp. gr. of about 2½, and consequently the strength of preparations will vary greatly, according to whether it is dispensed by weight or by measure.

IPECACUANHÆ RADIX.

IPECACUANHA ROOT

FR, IPECACUANHA ANNELE, GRE, BRECHWURZEL, ITAL, IPECACUANA, SPAN, IPECACUANA.

The dried Root of *Psychotria Ipecacuanha*.

The description of the Root given in the *B.P.* excludes the Carthagena variety, the *U.S.P.* includes both Rio and Carthagena.

Ipecacuanha, the *PG* only the Rio variety The Root official in the *BP* is not required to yield any definite percentage of alkaloids, that of the *USP* must contain not less than 1.75 p.c. of Ipecacuanha alkaloids, whilst that official in the *PG* is required to indicate at least 2.0 p.c. of total alkaloids

The *Brussels Conference* agreed that only the root bark should be powdered, rejecting the woody portion The powder should have an alkaloidal strength of 2 p.c. In the Brazilian Ipecacuanha Root the proportions of Emetine to Cephaeline are as 75 to 25, in the Carthagena Root as 45 to 55

Umney and Swinton record (*CD* '99, ii 203, 226, *PJ* '99, ii 89, 114, 123), the results of an examination of Johor Ipecacuanha. The total alkaloids amounted to 1.7, of which 1.21 p.c. represented Emetine, 0.39 p.c. Cephaeline, and 0.07 p.c. Psychotrine, the percentage proportion being 72.94 to 22.94 to 4.12. The results point to the conclusion that so far as the relative proportions of alkaloids are concerned, this root is practically identical with the Brazilian, but it contains a lower percentage of total alkaloids than the average Brazilian roots unmixed with stems

The relative percentage composition of the alkaloids from Brazilian and Columbian Ipecacuanha is given by Paul and Cownley (*JJP* '01, 115) as follows —Brazilian, Emetine 72.14, Cephaeline 25.87, Psychotrine 1.99 Columbian, Emetine 10.5, Cephaeline 56.8, Psychotrine 2.7 The paper gives an exhaustive *resumé* of the chemistry of Ipecacuanha

The active principle resides in the bark, the inner or woody part contains but little

From the experiments by Paul and Cownley (*PJ* (3) xxiv 61), it would appear (1) that the percentage of *total alkaloids* in Brazilian Ipecacuanha root does not vary much from 2 p.c. Rio Ipecacuanha Root contains the three alkaloids in the following proportions as compared with Carthagena and Indian Ipecacuanha —

Brazilian (root)—Emetine 1.45 p.c., Cephaeline 0.52 p.c., Psychotrine 0.04 p.c. Total 2.01 p.c.

Brazilian (stem)—Emetine 1.18 p.c., Cephaeline 0.59 p.c., Psychotrine 0.03 p.c. Total 1.80 p.c.

Columbian—Emetine 0.89 p.c., Cephaeline 1.25 p.c., Psychotrine 0.06 p.c. Total 2.20 p.c.

Indian—Emetine 1.79 p.c., Cephaeline 0.5 p.c., Psychotrine 0.09 p.c. Total 1.98 p.c. —Paul and Cownley, *PJ* '96, i 321, '02, ii 256

In 1893 it was stated by Paul (*PJ* (3) xxiv 212) that from so called de-emetinised Ipecacuanha he had obtained nearly 0.5 p.c. of the ordinary alkaloids of Ipecacuanha, but it can now be obtained entirely free from Emetine (*Pulvis Ipecacuanhae sine Emetina*)

Medicinal Properties.—Expectorant, diaphoretic, gastro-intestinal stimulant, cholagogue Emetic, slow in action (20 to 30 minutes), and depressant in large doses Used in emetic doses in whooping cough and croup to expel exudation on membrane as well as for its depressing effects on the circulation Used as an expectorant in acute and chronic bronchitis when the phlegm is thick and scanty, and in winter-cough and phthisis. Given in gouty dyspepsia and biliousness It relieves some forms of vomiting, such as that of pregnancy or alcoholism, when given in small doses, 1 or 2

minims of the **Vinum** every half-hour. Applied to the bites and stings of insects. The diaphoretic effect is best obtained when given in the form of the Compound Powder. In small doses it is commonly added to pills for chronic constipation. A **spray** of the Wine of Ipecacuanha has been strongly recommended by Ringer and Murrell for chronic bronchitis and asthma. Had been abandoned in many parts of the world in treatment of dysentery, but Manson has reintroduced it, with certain important improvements, in chronic dysentery of the anergic variety. Beginning with 30 grains, preceded by laudanum, and taken in diminishing dose every night for a week, is successful in most cases. For details see L '07, ii 1591.

In pneumonia — L '02, i 183

10 to 40 minims of the Wine three times a day in epilepsy. — L '98, ii 751; P J '99, i, 293

Dose — As an expectorant, $\frac{1}{2}$ to 2 grains = 0.016 to 0.13 gramme, as an emetic, 15 to 30 grains = 1 to 2 grammes.

Swiss, maximum single dose, 0.1 gramme = $1\frac{1}{2}$ grains; maximum daily dose, 0.5 gramme = 7 $\frac{1}{2}$ grains, maximum dose as an emetic, 5 grammes = 77 grains.

Prescribing Notes — Prescribed in small doses, *as a cathartic*, in the form of a powder, pill, cachet, or (in the form of a solution) can be made by using Dispensing Syrup, q s.

Tablets of Compound Ipecacuanha Powder may be obtained containing $\frac{1}{2}$, 1, 2, 3 and 5 grains. Simple Ipecacuanha, $\frac{1}{2}$, 1, 2, 3, 4 and 5 grains, De-emeticised, $\frac{1}{2}$, 1, 2, 3, 4 and 5 grains, Wine, 1, 2, 3, 4 and 5 minims; Ipecacuanha and Opium (B P pills), 4 and 5 grains.

Incompatibles — Lead and Mercury salts, vegetable Acids, and most Infusions.

Official Preparations — O the Root, Liquidum, Pulvis Ipecacuanhae, Proct. Ipecacuanhae, Morphin. et Ipecacuanhae, Liquid Extract, Acet. Ipecacuanhae, Vinum Ipecacuanhae, of the Compound Powder, Pilula Ipecacuanhae.

Not Official — Liqueur Ipecacuanhae, Tinctura Ipecacuanhae, Miscible, Glycerole of Ipecacuanha, Glycerin. Ipecacuanhae, Ipecacuanhae Mistura Ipecacuanhae Ammoniata, Mistura Ipecacuanhae Sacra, Mistura Ipecacuanhae cum Soda, Oxymel Ipecacuanhae, Pilula Ipecacuanhae, Symplic. Ipecacuanhae, Symplic. Ipecacuanhae Aceticus, Tinctura Ipecacuanhae, Tinctura Ipecacuanhae et Opium, Emetine, Emetine Hydrobromide, Emetine Hydrochloride, Vinum Emetine, Vinum Emetine Cephaeline, Cephaeline Hydrochloride, Psychotrine.

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S. Powder of Ipecacuanha in Austr, Belg, Dutch, Fr, Ger, Jap, Span, Swiss and U S contains 2 p c of alkaloids.

The Brussels Conference agreed that the powder should contain only the root-bark rejecting the woody portion, also that the powder should have an alkaloidal strength of 2 p c.

Descriptive Notes — In the B P. the name of the Ipecacuanha plant is given as *Psychotria Ipecacuanha*, Stokes, in the P G. as *Urugoga Ipecacuanha*, Baill. and in the U S P. as *Cephaelis Ipecacuanha*, A. Richardson. The name *Psychotria* is given on the ground that there is no good reason for rejecting the name *Psychotria*. The German name is *Wurde*.

priority, if kept distinct from *Psychotria Carthagenae* Ipecacuanha, official in the *USP* is stated to be derived from *Cephaelis acuminata*, Karsten

The Ipecacuanha Root of commerce is imported under the name of Rio, Matto Grosso, or Minas Gerais Ipecacuanha, from Brazil, where it is indigenous, and under the name of Johor or Selangor Ipecacuanha from the Federated Malay States, where it is cultivated. Ipecacuanha imported from the United States of Columbia, and known as Carthagenae Ipecacuanha, is official in the *USP* but not in the *PB* or *PG*. These kinds may be distinguished as follows.—The Brazilian Ipecacuanha is usually of a rusty dull brown colour but varies somewhat in tint, for a good deal of the Root arrives in a mouldy state, and after being washed it assumes a blackish-brown colour. The Root is about $\frac{1}{8}$ inch in diameter, and annulated or ringed to the extent of about 20 rings to the inch, much broken and rarely branched. It is officially described as occurring in tortuous pieces, not often exceeding 6 inches (15 cm) and $\frac{1}{4}$ inch (6 mm) in thickness, of a dark brick-red to very dark brown colour. The Selangor Root is similar, but as a rule there are more slender rootlets present and the root is branched. The bark is thick in proportion to the woody centre, horny and whitish in fracture (resinous but sometimes starchy, *BP*), and forms about 75 to 80 p.c. of the Root. Carthagenae Ipecacuanha is larger, of a paler and more distinctly reddish-brown tint, has more distant rings, which take the form of thinner merging ridges, and the fracture of the bark is greyish. Ipecacuanha Root varies much in quality, according to the amount of woody rhizome or stem present. The latter is smooth, slender, and cylindrical, with a very thin bark, and as the active principles are chiefly contained in the bark, only a very small quantity (about 5 p.c. of the amount present) being contained in the woody centre, the medicinal value of the Root is less in direct proportion to the amount of stem present. Brazilian and Selangor Ipecacuanha contain more Emetine than Cephaeline; Carthagenae Ipecacuanha contains more Cephaeline than Emetine.

In powder, Ipecacuanha is distinguished by the absence of vessels, sclerenchyma and bast fibres, and the presence of tacheids, porous parenchymatous cells and acicular raphides, and by starch grains which do not exceed 0.012 mm (*BP* (+)) and occur in groups of 3 to 5 grains. Several other roots have been offered as substitutes for Ipecacuanha, the histological distinctions of which are given in the *Pharm. Jour.* (3) xxxiv 210. Powdered Ipecacuanha Root has been found adulterated with almond meal, which may be recognised by the presence of the granular albuminous matter, some of which exhibits a crystalloid aspect, as seen under polarised light.

Tests.—Numerous methods have been suggested for the determination of the alkaloids in Ipecacuanha Root. The most complete research upon the chemistry of the alkaloids is undoubtedly that of Paul and Cownley. The method suggested by them (*A.J.P.*

'01, 116) is as follows — A weighed quantity of 50 grammes of the Root is mixed with one-fifth of its weight of Lime moistened with Water and extracted with Amyl Alcohol. The Amyl Alcohol solution is extracted with dilute acid, and the acid liquid shaken with Ether and Ammonia Solution to remove the Emetine and Cephaeline, leaving the Psycholine to be extracted from the ammoniacal liquid by Chloroform. The Ether residue is titrated with Semi-normal Volumetric Hydrochloric Acid Solution, 1 c.c. of the Volumetric Acid Solution representing 0.123175 gramme of Emetine and 0.11622 gramme of Cephaeline. A separation of the Emetine and Cephaeline is then effected by treating the Hydrochloric Acid solution with Sodium Hydroxide in the presence of Ether and repeatedly shaking the ethereal solution with Sodium Hydroxide Solution until all the Cephaeline has been separated. The Ether solution of the Emetine is evaporated and the residue titrated with standard acid, the result being expressed as Emetine. The Sodium Hydroxide solution, rendered alkaline with Ammonia Solution and shaken with Ether, the Ether residue of Cephaeline being titrated with Semi-normal Volumetric Acid Solution. The sum of the number of c.c. of Semi-normal Volumetric Hydrochloric Acid used in titrating the separated bases should equal the number required before their separation.

When a determination of the total alkaloidal content of a root is necessary, several good processes are available. That devised by Bird may be carried out and has the advantage that the difference between the volumetric and gravimetric determinations is reduced to a minimum. A weighed quantity of 10 grammes of the Ipecacuanha Root in fine powder is mixed with 1 gramme of Sodium Bicarbonate, and is then rubbed to a uniform moist granular powder with another 1 gramme of Sodium Bicarbonate shaken with 5 c.c. of Water. The moistened powder is then added to 20 c.c. of a mixture of 1 volume of Amyl Alcohol, 1 volume of Chloroform, and 3 volumes of Ether, contained in a separator, the stem of which is plugged with a pledget of cotton-wool, and the neck of which can be connected with a pressure bellows. The maceration is allowed to proceed for half an hour, with occasional shaking. The liquid is forced out of the separator by the pressure bellows, and the powder is again extracted with 10 c.c. of the same menstruum. After a vigorous agitation it is allowed to stand for 15 minutes, the liquid is again forced out with pressure bellows. The extraction is repeated ten or a dozen times at 15 minutes' intervals, with successive quantities of the same menstruum, or until the powder is exhausted. The ethereal liquids are mixed and extracted successively, first, with 10 c.c. of a mixture of 4 c.c. of Volumetric Sulphuric Acid Solution, and then with three successive quantities, each of 5 c.c. of Water, the aqueous layers being separated in each instance. The acid and aqueous solutions are mixed, rendered alkaline by the addition of 0.5 gramme of Ammonium Bicarbonate, and the liberated alkaloids are shaken out, first with 20 c.c., subsequently with two successive quantities, each of 10 c.c. of Chloroform, containing one-sixth its volume of Ether, and lastly with 20 c.c. of the same mixture plus 1 drop of Strong Ammonia.

the chloroformic solution being separated in each case, they are mixed and shaken with a saturated Sodium Chloride Solution containing 1 drop of Strong Ammonia Solution. The perfectly clear chloroformic solution is separated by forcing it through a very small plug of cotton-wool previously saturated with Chloroform and placed in the neck of the separator, the brine is washed by rotating it with a few cc more Chloroform mixture, the Chloroform solution separated, mixed with the first chloroformic solution, evaporated to about 1 or 2 cc, 5 cc of Ether sp gr 0.717 added, and the evaporation continued, allowing the residue to form a thin film on the inner surface of the vessel. Dry below 80° C (170° F) and weigh. The residue is dissolved in a neutral mixture of 10 cc Amyl Alcohol, 10 cc of Ether sp gr 0.717, and 5 cc of a saturated Sodium Chloride Solution, and titrated with Tenth-normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Methyl Orange Solution as the indicator of neutrality. 1 cc of Tenth-normal Volumetric Acid Solution may be taken as representing 0.021287 gramme of alkaloids. The factor 0.021287 is based upon the percentage composition of *Rao Ipecacuanha Root* recorded in the researches of Paul and Cownley, viz., practically 3 equivalents of Emetine to 1 of Cephaeline, using *B.P.* atomic weights. Where only approximate or comparative results are required, the following modification of the above process may be adopted. A weighed quantity of 12 grammes of the finely-powdered Root is mixed with 1 gramme of Sodium Bicarbonate, rubbed in a small mortar to a uniform moist granular powder with 1.4 gramme of Sodium Bicarbonate, shaken with 6 cc of Water, and added to 120 cc of a mixture of 1 volume of Amyl Alcohol, 1 volume of Chloroform, and 3 volumes of Ether, contained in a similarly-fitted separator to that used in the previous determination. The mixture is shaken occasionally during 2 or 3 hours, 5 cc. of brine added, agitated, and after aggregation of the powder 100 cc of the clear liquid, representing 10 grammes of the Root, is forced out by means of the pressure bellows into a graduated 100 cc flask. The alkaloids are then extracted by shaking with the acid mixture, and the process continued on the lines indicated above.

The method of determination adopted by the *U.S.P.* is essential as follows:—A weighed quantity of 15 grammes of the Root in No. 80 powder is shaken for 5 minutes in an Erlenmeyer flask, with a mixture of 115 cc of Ether and 35 cc of Chloroform, and, after the addition of 3 cc of Ammonia Water, is again shaken at intervals during half an hour. 10 cc of Water is added, and the liquid shaken until the powder agglomerates, when a measured quantity of 100 cc of the clear ethereal solution is transferred to a separator and the alkaloids extracted by shaking moderately for 2 minutes with a mixture of 10 cc of Normal Volumetric Sulphuric Acid Solution and 10 cc of Water. The acid aqueous liquid is separated, transferred to a second separator, and the extraction of the ethereal solution repeated, first with a mixture of 3 cc of Normal Volumetric Sulphuric Acid Solution and 5 cc of Water, and then with 10 cc of Water, the acid aqueous and the aqueous shakings being in each case separated and

transferred to the second separator. The mixed liquids in the second separator are now rendered alkaline by the addition of a sufficiency of Ammonia Water, and the alkaloids extracted by shaking for 1 minute, first with 25 c.c., then with 20 c.c., and lastly with 10 c.c. of Ether, separating the ethereal liquids in each case from the lower alkaline aqueous portion and transferring the ethereal solutions to a tared flask. The Ether is distilled off on a water-bath, and the residue is dissolved by gently warming it on a water-bath with 12 c.c. of Tenth-normal Volumetric Sulphuric Acid Solution, the excess of Volumetric Acid Solution being titrated with Fifteenth-normal Volumetric Potassium Hydroxide Solution, 5 drops of Cochineal Test Solution being used as an indicator of neutrality. The number of c.c. of Fifteenth-normal Volumetric Alkali Solution used is divided by 5, the quotient subtracted from 12, and the difference multiplied first by 0.0238, and then by 10, the result being the percentage yield of Ipecacuanha alkaloids present in the sample. It will be noticed that the factor employed by the *U.S.P.* in calculating the result of the Volumetric Test represents the mean combining weights of Emetine and Cephaeline. The choice of Ether as a solvent is open to question. Bird has shown that Ether alone is not entirely satisfactory as a solvent, as it is very difficult to remove the last traces of alkaloid from an alkaline solution by this reagent. Psychotrine is, moreover, according to Paul and Cownley, not extracted by Ether. The *P.G.* employs a mixture of 3 parts by weight of Ether and 1 part by weight of Chloroform for the extraction of the alkaloids, the method of determination being also a volumetric one. Sodium Hydroxide Solution (15 p.c.) is employed to liberate the alkaloids from their combinations. Upon the solubility of Cephaeline in Sodium Hydroxide Solution is based the method for its separation from Emetine, and its incomplete extraction by the Ether-Chloroform solvent therefore renders the results obtained by the process below the truth. The method of determination is briefly as follows.—A weighed quantity of 12 grammes of the Root in fine powder dried at 100° C (212° F.) is mixed in a well-stoppered bottle with 90 grammes of Ether and 30 grammes of Chloroform, and, after being well shaken 10 c.c. of a mixture of 2 parts by weight of Sodium Hydroxide Solution (15 p.c.) and 1 part by weight of Water is added, and the whole allowed to stand for 3 hours, with frequent intervals of vigorous shaking. 10 c.c. or sufficient Water to cause the powdered Root to agglomerate and the Ether-Chloroform solution to separate to a clear liquid is added, and, after being allowed to stand for 1 hour, a measured quantity of 100 grammes of the clear Ether-Chloroform solution is filtered through a dry, well-covered filter into a flask, and about one-half of the liquid is distilled. The residual liquid in the flask is transferred to a separator, the flask washed with three successive quantities, each of 5 c.c. of Ether, and the alkaloids thoroughly extracted with 12 c.c. of Tenth-normal Volumetric Hydrochloric Acid Solution. When the liquids have completely and clearly separated, and after the addition of sufficient Ether to cause the Ether-Chloroform solution to float on the surface of the acid liquid, the latter is filtered through a

small filter paper moistened with Water into a flask of 100 cc capacity. The Ether-Chloroform solution is shaken with three successive quantities, each of 10 cc of Water, which are separated and filtered through the same filter, the latter is washed with Water, and the mixed liquids are diluted with Water to 100 cc. A measured quantity of 50 cc is transferred to a white glass stoppered flask, having a capacity of 200 cc, mixed with 50 cc of Water and sufficient Ether to form a layer of about 1 cm. After the addition of 5 drops of a 1 in 500 solution of Iodeosin in Alcohol (90 p.c.), sufficient Hundredth-normal Volumetric Potassium Hydroxide Solution is added to cause the lower aqueous layer to assume a pale rose-red tint, not more than 20 cc of the Hundredth-normal Volumetric Solution should be necessary. The number of cc of Hundredth-normal Volumetric Potassium Hydroxide used, divided by 10, the quotient multiplied by 2, the product subtracted from 15, and the remainder, multiplied first by 0.024475, and then by 10, gives the percentage of Ipecacuanha alkaloids present in the sample, using the *German* atomic weights, and the proportions of 3 of Emetine and 1 of Cephaeline. If the mean molecular weights of Emetine and Cephaeline are employed, the factor is 0.024125. No factor is given in the *P.G.*, but if the *P.G.* official limit is calculated with the factor 0.024475, it represents not less than 1.96 p.c. In the process given by Keller, Ammonia Solution is employed for the liberation of the alkaloid, consequently there is not the same liability to the retention of Cephaeline. The detail of the method is as follows — A weighed quantity of 12 grammes of the powdered Root is shaken in a glass-stoppered bottle with 90 grammes of Ether and 30 grammes of Chloroform. After an interval, 10 cc of Ammonia Solution is added, the mixture shaken at intervals for half an hour, and, after the addition of 10 cc of Water, it is again shaken for a short time until the powder agglomerates. The clear Ether-Chloroform solution is filtered through a small filter paper, which has been moistened with Ether, and a weighed quantity of 100 grammes is introduced into a separator and shaken with three successive quantities, each of 25, 15, and 10 cc of a 1 p.c. Hydrochloric Acid Solution, the shakings being repeated if necessary so long as a few drops of the acid shakings yield a precipitate with Potassio-Mercuric Iodide (Mayer's) Solution. The acid aqueous solutions are separated, mixed, transferred to a separator, made alkaline with Ammonia Solution, and shaken with three successive quantities, each of 50, 30, and 20 cc, of a mixture of 2 parts by weight of Ether to 3 parts by weight of Chloroform. The Ether-Chloroform solutions are separated in each case, mixed, filtered through a small filter paper moistened with Ether, into a tared flask, the Ether-Chloroform distilled, the residue dried on the water-bath till constant in weight, cooled and weighed. In the event of the titration of the alkaloidal residue being required, it may be dissolved in 5 cc of neutral Alcohol (90 p.c.), Water added until the liquid becomes faintly opalescent, and the titration effected with Tenth-normal Volumetric Hydrochloric Acid Solution, using Hæmatoxylin Solution (1 in 500 of Alcohol, 90 p.c.)

as an indicator Keller states that each c.c. of Volumetric Acid is equivalent to 0.0254 gramme of alkaloids, which agrees with the $C_{30}H_{40}N_2O_5 \cdot 2HCl$ formula of neutral Emetine Hydrochloride. If the 3:1 ratio of Emetine to Cephaeline factor is used, each c.c. of Tenth-normal Volumetric Acid will be equivalent to 0.024287 gramme of Ipecacuanha alkaloids.

Allen has studied the comparative colour reactions of the Ipecacuanha alkaloids and the Opium alkaloids (see also Tinctura Camphoræ Composita), and the results of his experiments are recorded (*Analyst*, '02, 349). The colour tests were made by taking up the alkaloidal solution in a pipette and allowing it to fall drop by drop on to the concave side of a porcelain crucible lid placed on a flask full of boiling Water. To the spot of alkaloidal residue thus obtained a drop of reagent was added by means of a glass rod, and the mixture cautiously stirred. With Ferric Chloride the Ipecacuanha alkaloids gave a blue coloration changing to green, as against a greenish-blue from a residue of Opium alkaloids, Sulphuric Acid containing 0.5 p.c. of Molybdic Acid (Fehde's reagent) gave colours varying from bluish-purple to violet, the colour resembling that given by Opium alkaloids, but not so bright as that yielded by pure Morphine, Starch and Iodic Acid gave, with some specimens of alkaloidal residue, an immediate blue coloration, but a negative or hardly result in other cases, Opium alkaloids gave an immediate blue colour, with Ferric Chloride and Potassium Ferricyanide, both the Ipecacuanha alkaloids and the Opium alkaloids gave an immediate Prussian blue coloration. In some cases the alkaloidal residues were purified by treatment with Lead Acetate. The isolated and purified alkaloids in some instances gave reactions less striking than those obtained with the mixed alkaloidal residue.

For a valuable means of detecting Ipecacuanha alkaloids, see Psychotrine, p. 693.

A determination of the total ash of powdered Ipecacuanha is stated (*Proc Amer Jour Pharm* 51, 771, *PJ* '03, 357) to afford but little clue to the nature of the drug powdered. The presence of more than 1 p.c. of sand generally indicates a dusty or otherwise objectionable root.

Preparations.

ACETUM IPECACUANHÆ VINEGAR OF IPECACUANHA

Liquid Extract of Ipecacuanha, 1, Alcohol (90 p.c.), 2, Diluted Acetic Acid, q.s. to make, 20 (1 in 20)

Dose — 10 to 30 minims = 0.6 to 1.8 c.c.

Tests — Vinegar of Ipecacuanha has a sp. gr. of about 0.990; contains about 0.75 p.c. w/v of total solids, about 12.0 p.c. w/v of Absolute Alcohol, and about 3.57 p.c. w/v of absolute Acetic Acid. A measured quantity of 10 c.c. requires about 6 c.c. of Normal Volumetric Sodium Hydroxide Solution for neutralisation, Phenolphthalein Solution being employed as an indicator of neutrality, this corresponding to about 3.57 p.c. w/v of absolute Acetic Acid. It is prepared with the official Fluid Extract, which is required to

contain not less than 2 and not more than 2.25 p.c. w/v of alkaloids, the Vinegar should therefore contain not less than 0.10 p.c. w/v nor more than 0.112 p.c. w/v. No official method of determination is given, but a convenient process has been suggested by Bird. A measured quantity of 100 c.c. is nearly neutralised with Potassium Hydroxide Solution, 2.5 c.c. of Lead Subacetate Solution added, and a few grains of washed Asbestos powder. The mixture is heated on the water-bath until a distinct separation of the precipitate is observed, transferred to a Buchner filter, and filtered under pressure. The nearly dry solid cake remaining on the filter is washed with 30 c.c. of Water, added in small portions at a time, the filtrate and washings are mixed with 25 c.c. of diluted Sulphuric Acid, and the precipitated Lead Sulphate separated by again filtering through a Buchner's filter, and washed with 15 c.c. of Water. 25 c.c. of Chloroform is added to the mixed filtrate and washings, and Ammonia Solution in excess. The flask is corked, agitated vigorously, and the contents transferred to a separator, which is plugged at the neck with a pledget of Cotton-Wool. The chloroformic solution and a portion of the aqueous liquid is forced out of the first separator into a second separator, by means of a pressure ball attached to the former, and the clear chloroformic liquid is separated and transferred to a tared basin or flask. The chloroformic extraction is repeated with four successive quantities, each of 25 c.c. of Chloroform, the aqueous liquid in the second separator being in each instance returned to the first separator previous to the addition of the Chloroform. The mixed chloroformic solutions are evaporated to dryness on a water bath, the residue dried at a temperature below 80° C (176° F) and weighed.

EXTRACTUM IPECACUANHÆ LIQUIDUM LIQUID EXTRACT OF IPECACUANHA

A Liquid Extract standardised to contain 2 to 2½ grams of Ipecacuanha alkaloids in 110 minims (2 to 2.25 grammes in 100 c.c.); prepared with Ipecacuanha Root, Calcium Hydroxide, and Alcohol (90 p.c.)

Dose—As an expectorant, ½ to 2 minims = 0.03 to 0.12 c.c., as an emetic, 15 to 20 minims = 0.9 to 1.2 c.c.

Swiss, maximum single dose, 0.05 gramme, maximum daily dose, 0.25 gramme

Foreign Pharmacopœias—Official in Dan, Swed, Swiss, and U.S., 1 in 1 Belg (Ipecacuanha Extractum Fluidum Compositum) 3 of Tincture in 10. Fr, Mex and Span have solid extract. Swiss contains at least 2 p.c. of alkaloids and U.S. 1.75 p.c.

Tests.—Liquid Extract of Ipecacuanha has a sp. gr. of 0.885 to 0.910, contains from 6 to 12 p.c. w/v of total solids and about 78 p.c. w/v of Absolute Alcohol. The Fluid Extract official in the *B.P.* is required to contain not less than 2.0 p.c. w/v and not more than 2.25 p.c. w/v of alkaloids. The *B.P.* states 'alkaloid,' but, inasmuch as it refers immediately above to 'alkaloids,' this may be taken as a printer's error. The essential features of the official method of assay are as follows.—A measured quantity of 20 c.c. of the Fluid Extract is mixed with 20 c.c. of Water and evaporated on a

water-bath until the Alcohol is dissipated. The warm solution is precipitated with Lead Subacetate Solution, and is preferably allowed to remain on the water-bath until the precipitate becomes granular and commences to subside. It is then filtered off, washed with Water, and to the filtrate and washings sufficient Sulphuric Acid is added to precipitate the excess of Lead Subacetate. The precipitate is filtered off, washed with Water, and the mixed filtrate and washings are transferred to a separator. Sufficient Ammonia Solution to form an excess is added, and the liberated alkaloids are extracted by shaking with three successive quantities each of 25 c.c. of Chloroform. The chloroformic layers are removed in each case, mixed, transferred to a tared flask, evaporated on a water-bath, and the residue dried at a temperature below 80°C (176°F .) till constant in weight, and weighed. This weight multiplied by 5 yields the p.c. w/v of total alkaloids present in the sample. This process has been subjected to severer and more adverse criticism than almost any other assay process in the British Pharmacopoeia. The quantity of the Liquid Extract taken for the determination is too large and too wasteful of material. The Lead precipitate is bulky and unmanageable. Its nature renders the washing with Water a lengthy and troublesome operation. The chloroformic liquids have a tendency to obstinately emulsify, and they require a considerable time to separate, and over and above the inaccuracy due to the consequent unavoidable loss of alkaloid is the fact that the alkaloidal residue, when it is finally obtained, is impure.

Several processes of determination have been suggested with a view to devising an expeditious, accurate and readily-conducted method. The process devised by Wilson gives more accurate results than the *B.P.* method, and can, moreover, be almost completed whilst the first *B.P.* Lead precipitate is being filtered and washed. The method has been in general use in the author's laboratory, and has been found expeditious and accurate. Liquid Extracts which contain an exceptional amount of resinous substances are not suitable for assay by this process. In carrying out the process a measured quantity of 20 c.c. of the Liquid Extract is placed in a porcelain basin, diluted with 20 c.c. of Water, evaporated to somewhat less than half its bulk, and allowed to cool. It is mixed with 1 c.c. of dilute Sulphuric Acid, transferred to a separator, the dish washed with 20 c.c. of Water and the mixed liquids shaken with 10 c.c. of a mixture of equal parts of Chloroform and Ether, separation being promoted by gently warming, the Ether-Chloroform layer is separated and the shaking repeated with two successive quantities each of 10 c.c. of a similar mixture, the Ether-Chloroform layers being in each case separated and rejected. An excess of Ammonia Solution is now added and the liberated alkaloids are removed by shaking with 10 c.c. of Ether-Chloroform (equal volumes), the mixture warmed to promote separation, the Ether-Chloroform layer separated, and transferred to a tared flask. The extraction with Ether-Chloroform is repeated with two successive portions each of 10 c.c. of the Ether-Chloroform, the Ether-Chloroform layers being separated in each case as previously, and

transferred to the tared flask. The Ether-Chloroform is removed by distillation, the residue is dried at a temperature below 80°C (176°F) until constant in weight, cooled and weighed. It may then be dissolved in a measured quantity of Tenth-normal Volumetric Hydrochloric Acid Solution and the excess of acid titrated with Tenth normal Volumetric Sodium Hydroxide Solution, using a few drops of a 1 in 500 Iodeosin Solution in Alcohol (90 p c) as an indicator of neutrality. The factor to be used in calculating the result of the volumetric determination is discussed under the tests for Ipecacuanha Root. Naylor, in his Presidential address to the British Pharmaceutical Conference on the Standardisation of Galenicals, claims that the process detailed by Naylor and Bryant (*Y B P* '99, 345) and that of Farr and Wight (*Y B P* '99, 340) are the only published methods of assay which in his hands have yielded uniformly accurate results with every type of Liquid Extract. The process which is described in the *Y B P* '99, 345, consists in warming a measured quantity of 10 c.c. of the Liquid Extract in a basin over a water-bath until the Alcohol is dissipated. The residue is transferred to a 50 c.c. flask, the basin rinsed out with successive portions of a mixture of 2 c.c. of Diluted Sulphuric Acid and 30 c.c. of Water. The solution is filtered, and the filter washed with Water until the volume of the filtrate amounts to 50 c.c. A measured quantity of 25 c.c. (= 5 c.c. of the Liquid Extract) is transferred to a separator, the measure washed with Water, and the mixed liquids are shaken with 10 c.c. of Chloroform. The Chloroform layer is separated and rejected, the shaking being repeated with a further quantity of 10 c.c. of Chloroform, and the chloroformic layer again separated and rejected. The aqueous portion is then rendered alkaline with Ammonia Solution and the liberated alkaloids extracted by agitation with three successive quantities each of 10 c.c. of Chloroform. The chloroformic layer is in each case separated, transferred to a tared flask, the mixed chloroformic solutions evaporated to dryness, the residue dried till constant in weight, cooled and weighed. It may then be dissolved in a measured quantity of Tenth-normal Volumetric Hydrochloric Acid Solution and the excess of Volumetric Acid titrated as described above.

Bnd is rightly of opinion that whatever the inaccuracies and imperfections of the Pharmacopœia process, the fact remains that it is the standard by which the Liquid Extract must be tested and judged, and in the event of another process being selected it is absolutely essential that the results obtained shall bear a constant relation to the result given by the *B P* method and be capable of accurate translation into the official figures. He has devised a process, the details of which are claimed to be quite in accordance with a close interpretation of the *B P*. A measured quantity of 20 c.c. of the Liquid Extract is mixed with 20 c.c. of Distilled Water and a sufficient quantity of Acetic Acid to ensure a faintly acid reaction. The Alcohol is evaporated off, and 20 c.c. of Water and 10 c.c. of Liquor Plumbi Subacetatis added. The mixture is allowed to remain on the water-bath until the magma which first forms changes to a thin

liquid, and the precipitate assumes a finely granular condition. It is then transferred to a Buchner's funnel and filtered under pressure. The nearly dry solid cake remaining on the filter is washed with 30 c.c. of water added in successive small portions, 25 c.c. of diluted Sulphuric Acid is added to the filtrate and washings, and the precipitated Lead Sulphate removed by filtration on the Buchner's filter under pressure. The cake of Lead Sulphate is washed with 15 c.c. of Water, and to the filtrate and washings 5 c.c. of Chloroform and an excess of Ammonia Solution are added. The flask is cooled, the whole vigorously shaken, and the contents transferred to a separator, the neck of which is plugged with a pledget of Cotton-Wool, and the orifice of which is fitted with a cork and rubber bellows. The Chloroform layer and a portion of the aqueous layer is then forced out of the first separator into a second separator, from which the clear chloroformic layer is drawn off into a tared flask. The aqueous portion of the liquid in the second separator is transferred to the first separator, and the contents of the latter are shaken with two successive quantities, each of 25 c.c. of Chloroform, the chloroformic layers in each case being removed and transferred to the tared flask. The Chloroform is evaporated on a water-bath, the residue dried below 80°C (176°C) till constant. The alkaloidal residue is then dissolved in a neutral mixture of 10 c.c. Amyl Alcohol, 10 c.c. of Ether, and 5 c.c. of a saturated Sodium Chloride Solution, and titrated with Tenth-normal Volumetric Acid Solution, using Methyl Orange or Hematoxylin Solution as the indicator. The same factor as above mentioned may be employed in calculating the result of the volumetric determination.

The standard originally given for the Root was not less than 2 p.c. of Ipecacuanha alkaloids, whilst that for the Fluid Extract was 1.75 p.c. w/v, the present standard for the Root is 1.75 p.c., for the Fluidextractum 1.5 p.c.

The method adopted by the *USP* for the determination of the alkaloids is a volumetric one, the essential details being as follows — The Alcohol is removed from a measured quantity of 10 c.c. of the Fluid Extract by evaporation in a porcelain evaporating basin on a water-bath, 5 c.c. of Normal Volumetric Sulphuric Acid Solution and 20 c.c. of Water being added when almost cool, and the liquid stirred intermittently for 3 minutes. It is filtered into a separator, the dish and filter paper are washed with successive quantities of 10 c.c. and 5 c.c. of Water, the filtrate and washings are rendered alkaline with Ammonia Solution and shaken for 1 minute with 20 c.c. of Ether. The aqueous layer is separated and the extraction of the alkaloids repeated with two successive quantities each of 10 c.c. of Ether, the ethereal solutions being in each case separated and added to the first ethereal solution. The Ether is evaporated at a gentle heat from the mixed ethereal liquids and the alkaloidal residue is dissolved in 10 c.c. of Tenth-normal Volumetric Sulphuric Acid Solution. When completely dissolved 5 drops of Cochineal Solution are added, and the excess of Volumetric Acid Solution is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution. The number of c.c.

required is divided by 5, the quotient subtracted from 10, the remainder multiplied first by 0.0238, and then by 10, yields the percentage w/v of Ipecacuanha alkaloids present in the sample. The process has been tried in the author's laboratory, and works very satisfactorily. The separations are clean and sharp, the extracted alkaloids are of good colour, and in the volumetric determination no difficulty exists in determining the end-reaction. Ether being employed as a solvent, the residuo consists naturally of Ether soluble alkaloids only, Psychotrine being insoluble in Ether is not determined. In the case of an official preparation of Rio Ipecacuanha, according to Paul and Cownley the Psychotrine is hardly worth consideration. The process being a volumetric one yields lower results than are obtained by a gravimetric process, but returns the true alkaloidal percentage and not a percentage of alkaloids plus an indefinite amount of inert matter. A sample of *B.P.* Liquid Extract which yielded when examined by the *B.P.* process 2.0 p.c. w/v of total alkaloids, yielded when examined by the *U.S.P.* process 1.67 p.c. w/v of the alkaloids of Ipecacuanha.

The use of Ether, first, in a well diluted decidedly acid solution of the Liquid Extract, and then in ammoniacal solution, was suggested by Paul and Cownley, *Y.B.P.* '03, 100.

The Liquid Extract has been stated to deteriorate rapidly in alkaloidal content on keeping. Specimens of Liquid Extract prepared with both Rio and Carthagena Root are stated (*P.J.* '99, ii 622) to have deteriorated in two months from 2.08 p.c. to 1.528 p.c., indicating a loss of 26.53 p.c. in the case of the Liquid Extract made from the Rio Root, and from 2.1 p.c. to 1.525 p.c. in the case of the Liquid Extract made from the Carthagena Root, the Wine made from the Rio Root from 0.10 p.c. to 0.025 p.c. Deterioration is also stated (*P.J.* '00, i 8) to take place in the alkaloidal value of the Liquid Extract and more rapidly in the case of the Wine. On the contrary it has been stated (*P.J.* '99, ii 633, '00, i 54) that there is no reason for such depreciation, and it is shown that samples examined after the lapse of considerable intervals of time showed no depreciation in alkaloidal value. Such a marked depreciation as 26.53 p.c. in alkaloidal strength in two months is regarded (*P.J.* '00, i 414) as still awaiting explanation, but will probably be found due to the presence of much free alkaloid in alkaline or insufficiently acid solution. From the result of an examination of samples of the Liquid Extract (*C.D.* '02, ii 290, *P.J.* '02, ii 134) it appears that the total amount of alkaloids lost during nine months amounted to 5.66 p.c., so that although a depreciation occurs, the amount is very small. The indications are distinctly in favour of loss of alkaloid by precipitation as opposed to loss of alkaloid by decomposition.

PILULA IPECACUANHÆ CUM SCILLA. PILL OF IPECACUANHA WITH SQUILL

Compound Powder of Ipecacuanha, 3, Squill, in powder, 1; Ammoniacum, in powder, 1, Syrup of Glucose, *q.s.*

(about 1 of Opium in 20)

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

Foreign Pharmacopœias—Official in Port, similar to Brit. Not in the others

A corresponding *Pilula Ipecacuanhæ cum Urginea*, is Official in the *Ind* and *Coll*, and the Eastern Colonies

PULVIS IPECACUANHÆ COMPOSITUS. COMPOUND POWDER OF IPECACUANHA *B.P. Syn*—DOVER'S POWDER

Ipecacuanha Root, 1, Opium, 1, Potassium Sulphate, 8 All in powder (1 Opium, 1 Ipecac in 10)

Medicinal Properties—An admirable diaphoretic and anodyne, it is also most useful in vomiting, dysentery and diarrhoea, in the latter case 3 or 4 grains it will often relieve heartburn

Dose—5 to 15 grains = 0.3 to 1 gramme

Ph. Ger maximum single dose, 1.5 grammes, maximum daily dose, 5.0 grammes

Foreign Pharmacopœias—Official in all, and is the well-known Dover's Powder, Austr, Belg, Ger, Norw, Russ, Swed and Swiss (*Pulvis Ipecacuanhæ Opiatus*), Hung (*Pulvis Doveri*), Dan (*Pulvis Ipecac Thebaicus*), Dutch (*Pulvis Opii Compositus*), Fr (*Poudre d'Ipecacuanha Opiacée*), Port (*Po de Ipecac Composto*), Jap (*Pulvis Doveri*), and US (*Pulvis Ipecacuanhæ et Opii*), with Milk Sugar, all same strength as Brit, Span (*Polvere Ipecacuanha Opiado*), 1 Opium, 1 Ipecacuanha, in 10, Ital (*Polvere del Dover*), Opium 1, Potassium Sulphate 2, Mex (*Pulvis Ipecacuanhæ et Opii*), Opium 1 Ipecacuanha 1, Nitre 4, Potassium Sulphate 4

The *Brussels Conference* agreed for it to contain 10 p.c. of *Pulvis Opi*

The original Powder of Dr Dover was prepared by fusing together 4 parts of Potassium Nitrate with 4 of Potassium Sulphate, and reducing the product to fine powder: 1 of Ipecacuanha, 1 of Opium and 1 of Liqueur

TROCHISCUS IPECACUANHÆ. IPECACUANHA LOZENGE

$\frac{1}{4}$ grain of Ipecacuanha in each, with Fruit Basis

Dose—1 to 3 lozenges

Foreign Pharmacopœias—Official in Austr, Belg (*Tabellæ*), Dutch, Fr, Jap, Mex, Port, Russ, and Swiss, 0.01 gramme = about $\frac{1}{4}$ grain, Ital, 0.015 gramme = about $\frac{1}{4}$ grain; Not in the others

TROCHISCUS MORPHINÆ ET IPECACUANHÆ. See p 787

VINUM IPECACUANHÆ. IPECACUANHA WINE

Liquid Extract of Ipecacuanha, 1, Sherry, 19 (1 in 20)

Dose.—As an expectorant, 10 to 30 minims = 0.6 to 1.8 cc.; as an emetic, 4 to 6 fl drms = 14.2 to 21.4 cc

Foreign Pharmacopœias—Official in Swed, Fluid Extract 1, Marsala 10, US, Fluid Extract 1, Alcohol (95 p.c.) 1, White Wine 8, Dutch, Ipecacuanha 1, Diluted Spirit 1, Vinegar 9 (or), Jap, Norw and Russ, 1 of Ipecacuanha and 10 of Sherry, Port, 1 of Ipecacuanha in 20 of Port. Not in the others

Tests.—Ipecacuanha Wine has a sp gr of 0.985 to 1.000; contains about 5 p.c. w/v of total solids and about 20 p.c. w/v of Absolute Alcohol. It is prepared with the official Liquid Extract, which is required to contain not less than 2.0 p.c. w/v nor more than 2.25 p.c. w/v of total alkaloids. The Wine should, therefore,

contain not less than 0.1 g w/v nor more than 0.125 g w/v of total alkaloids. No official method is given for the determination of the alkaloids. A convenient method is that suggested by Bird. A measured quantity of the Wine is mixed with 2.5 cc of Lead Subacetate Solution and a few grains of washed asbestos powder. The mixture is heated on a water-bath until a distinct separation of the precipitate occurs and is then filtered under pressure on a Buchner's filter, the process being completed as described in Bird's process for the assay of the Liquid Extract (*Y B P '99*, 347).

The following process has been suggested by Naylor and Bryant. — A measured quantity of 100 cc is evaporated to one tenth its volume, a little Kieselsolium stirred in, the mixture transferred to a beaker, and the basin washed with a mixture of 2 cc of diluted Sulphuric Acid and 30 cc of Water. The solution is then filtered, and Water passed through the filter until the volume measures 50 cc. A measured quantity of 25 cc of the filtrate (= 50 cc of the Wine) is transferred to a separator and treated as described in Naylor's method for the assay of the Liquid Extract.

Not Official

ELIXIR IPECACUANHÆ — Liquid Extract of Ipecacuanha, 1, Rectified Spirit, 1, Simple Elixir, 1, Glycerin, 5, Water, *q s* to produce 20 — *Pharm Form*

This has been incorporated in the *B P C*

EXTRACTUM IPECACUANHÆ LIQUIDUM (MISCIBILE) — Liquid Extract of Ipecacuanha, *B P*, 100, Distilled Water, *q s* to produce 100 of finished product, Acetic Acid *q s*

Mix the Liquid Extract of Ipecacuanha with 100 of Distilled Water, allow to stand in a cool place for 24 hours, filter, washing the residue on the filter paper until colourless, keeping the washings separate, acidify the filtrate with Acetic Acid to a slight acid reaction. Distil on a water-bath until the distillate (as shown by volume and *sp gr*) contains 45 absolute Alcohol. This will generally measure about 52. Reserve this portion of the distillate, continue distillation to recover remaining Alcohol. Evaporate the residue on the water bath to about 42, allow to cool, pour off the bright liquid, add this to the reserve distillate. Rinse the dish with the washings obtained in the first part of the process, filter, and evaporate to make the total volume 100. — *F C J Bird, C D '99*, n 220

This has been incorporated in the *B P C*

GLYCEROLE OF IPECACUANHA — Liquid Extract of Ipecacuanha, 100, Distilled Water, 100. Mix as in Miscible Extract, allow to stand, filter, wash the residue, evaporate the washings separately, acidify the filtrate with Acetic Acid to a very faint acid reaction, distil off the Alcohol and evaporate on a water bath (adding the evaporated washings towards the end) to 50, add Glycerin, 50. (1 in 1)

This forms a clear solution with detannated wine, syrups, or aqueous liquids. It contains the *B P* proportion of alkaloid, being the same strength as the Liquid Extract, and for many obvious purposes furnishes a convenient preparation of Ipecacuanha. — *F C J Bird, C D '99*, n 220

Glycerinum Ipecacuanhæ *Syn* Glycerol Ipecacuanhæ, — Vinegar of Ipecacuanha, 50, Glycerin, 50 — *B P C*

This contains 24 of Liquid Extract in 100, and is, therefore, only $\frac{1}{4}$ of the strength of the above preparation.

LINCTUS IPECACUANHÆ — Vinegar of Ipecacuanha, Syrup of Tolu, Glycerin, Mucilage of Tragacanth, of each equal parts — *St Thomas's*

This has been incorporated in the *B P C*

MISTURA SCILLÆ ET IPECACUANHÆ (see p. 1065).

MISTURA IPECACUANHÆ AMMONIATA—Ipecacuanha Wine, 10 minims, Ammonium Carbonate, 3 grains, Peppermint Water, to 1 fl oz—*St Mary's*

MISTURA IPECACUANHÆ SALINA—Ipecacuanha Wine, 6 minims, Spirit of Nitrous Ether, 20 minims, Paregoric Elixir, 20 minims, Solution of Ammonium Acetate, 1 drim, Water, to 1 fl oz—*St Mary's*

MISTURA IPECACUANHÆ CUM SODA—Sodium Bicarbonate, 15 grains, Ipecacuanha Wine, 10 minims, Aromatic Spirit of Ammonia, 15 minims, Peppermint Water, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

OXYMEL IPECACUANHÆ—Liquid Extract of Ipecacuanha, 2 50, Oxymel, *qs* to produce 100—*B P C*

PILULA IPECACUANHÆ CUM URGINEA.—Compound Powder of Ipecacuanha, 3, Urginea, dried and in powder, 1, *in powder*, 1, Syrup of Glucose *qs* Dose—4 to 8 grains This Pill contains about 5 p.c. of Opium—*Ind and Col Add*

This has been incorporated in the *B P C*

SYRUPUS IPECACUANHÆ ACETICUS.—Vinegar of Ipecacuanha, 20 fl. oz, Refined Sugar, 86 oz—*B P C Formulary 1901*, incorporated in the *B P C* under the title of **Syrupus Ipecacuanhæ** Altered in the *B P C Supplement* to 75 or refined sugar dissolved in 50 of Vinegar of Ipecacuanha, adding sufficient Distilled Water to produce 100

Dose—15 to 120 minims = 0 9 to 7 1 cc

Syrupus Ipecacuanhæ—Tincture of Ipecacuanha, 1, Simple Syrup, 100—*Austr, Dutch, Jap, Span and Swiss*

Belg—Tincture of Ipecacuanha, 1, Simple Syrup, 10 (evaporated to 10) also Ipecacuanha Syrupus Compositus Compound Liquid Extract, 1, Simple Syrup, 20 U S—Fluid Extract of Ipecacuanha, 7, Acetic Acid, 1, Glycine, 10, Water, to 100

Ger—Ipecacuanha, 1, Alcohol (90 p.c.), 5, Water, 40, digest 48 hours, filter, and to make 100 add 60 of Sugar, and dissolve to make 100 of Syrup

Fr—Alcoholic Extract of Ipecacuanha, 1, Alcohol (70°), 3, Simple Syrup 100 *Fr* has also a Compound Syrup

Mex—Ipecacuanha, 1, Alcohol (60 p.c.), 4, Simple Syrup, 95

All by weight except U S

The *Brussels Conference* agreed that the Syrup should be prepared with 10 p.c. of the tincture

TINCTURA IPECACUANHÆ—Bruised Ipecacuanha, 1, Alcohol (60 p.c.), 10

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Span, and Swiss, 1 in 10, Jap, 1 and 10, Hung, Mex and Port, 1 in 5 All by weight Not in the others

Austr, Belg, Jap and Swiss at least 0 2 p.c. of alkaloids

The *Brussels Conference* agreed to a strength of 10 p.c. for the Tincture prepared by percolation with Alcohol (70 p.c.)

TINCTURA IPECACUANHÆ ET OPII *Syn* Fluid Dover's Powder—Tincture of Deodorised Opium, 100 Fluid Extract of Ipecac, 10, Alcohol (49 p.c.), *qs* to produce 100 Evaporate the Tincture of Deodorised Opium in a tared dish on a water-bath until it weighs 80 Water-bath Extract of Ipecac, filter, and pass enough Alcohol (49 p.c.) over the filter to make 100 Average Dose—8 minims (0 50 cc)—*U S P*

This has been incorporated in the *B P C*, employing Extractum Ipecacuanhæ Liquidum, *B P*, and Alcohol (60 p.c.)

EMETINA $C_{15}H_{21}NO_7$, eq 245 85 or $C_{15}H_{21}N_2O_4$, 490 70—A colourless amorphous base present in varying amount in Brazilian, Columbian and Indian Ipecacuanha Root, as given under Ipecacuanha. On exposure to light it rapidly

acquires a yellow colour. It is readily soluble in Alcohol, Ether, Chloroform and Benzene, but sparingly in Water.

The chief salts for medicinal purposes are the Hydrochloride and Hydrobromide.

The name 'Emetine' used to be applied to an impure extractive, containing the mixed alkaloids of *Ipecacuanha*, which is now listed as **Emetine (impure)** or **Emetine (extract)**.

Tests—Emetine melts at about 68°C ($154\frac{1}{2}^{\circ}\text{F}$). It is strongly alkaline in reaction towards Litmus, and neutralises acids completely, forming salts which are neutral in reaction towards the ordinary indicators of neutrality. It may therefore be readily determined by titration with Normal Volumetric Hydrochloric Acid Solution, using either Methyl Orange Solution or Iodocyan Solution as an indicator. 1 cc of Normal Volumetric Hydrochloric Acid Solution is equivalent to 0.24535 gramme of Emetine. When precipitated from the solution of one of its salts by Potassium or Sodium Hydroxide Solution, Emetine is insoluble in an excess of the reagent. This test distinguishes Emetine from Cephaeline. When dissolved in sufficient Hydrochloric Acid to effect solution and to show a slight excess, it yields, with Platinum Chloride Solution, a buff coloured amorphous precipitate, almost insoluble in Water or Alcohol. According to Allen (*Analyst*, vol. 349), it yields with Ferric Chloride an indefinite colour reaction, with Sulphuric Acid containing 0.5 per cent w/v of Molybdic Acid (Fehde's reagent), a dirty green coloration, if obtained from *Rio Ipecacuanha*, and a bluish coloration if obtained from Carthagena root, with Fehde's reagent and Hydrochloric Acid a grass green coloration, with Starch and Iodic Acid, a negative reaction, and with Ferric Chloride and Potassium Ferricyanide a gradual blue coloration. It should leave no residue when ignited with free access of air.

EMETINÆ HYDROBROMIDUM $\text{C}_7\text{H}_7\text{NO}$, HBr , $2\text{H}_2\text{O}$, eq 362.46.—Crystallises from Water in beautiful silky tufts of needles. Although readily soluble in Water, it is much less soluble than the Hydrochloride, difficultly so in Absolute Alcohol or in Chloroform. The commercial salt contains 67.95 per cent of alkaloid. It is rendered anhydrous at 100°C (212°F).

Tests—Emetine Hydrobromide dissolves in Water. The solution yields on the addition of Potassium or Sodium Hydroxide Solution a precipitate insoluble in excess of the reagent. It yields when faintly acidified with Nitric Acid a yellowish-white curdy precipitate, which when washed is almost insoluble in Ammonia Water and in Nitric Acid, but is readily dissolved by Potassium Cyanide Solution, on the cautious addition of a drop or two of Chlorine Water to the aqueous solution, it yields a yellowish coloration, and if the liquid be shaken with a few cc of Chloroform the yellowish coloration passes into the chloroformic layer. The separated alkaloid should answer the tests distinctive of Emetine given under Emetina. It should leave no residue when ignited with free access of air.

EMETINÆ HYDROCHLORIDUM—Crystallises from Water in radiating groups of silky filaments. Very soluble in Water, and in Alcohol. Dried at 100°C the salt is rendered anhydrous, and then has the composition $\text{C}_7\text{H}_7\text{NO}_2\cdot\text{HCl}$, eq 282.51. When crystallised from an acid solution $\text{C}_7\text{H}_7\text{NO}_2\cdot\text{HCl}$, $3\text{H}_2\text{O}$, eq 336.18. Both salts are permanent, undergoing no alteration in colour after being kept for some months.

Medicinal Properties—A powerful emetic and expectorant. For all the uses of *Ipecacuanha* where vomiting is not desired, Emetine in small doses seems likely to prove of considerable value, also as an emetic in larger doses of from $\frac{1}{2}$ to 1 grain when a more depressing action is required. The powerful local constricting effect upon blood vessels may also prove useful in hyperæmic and inflammatory conditions. The emetic dose of Emetine is about double that of Cephaeline. Emetine caused a flow of watery mucus from the nasal mucous membrane when a full dose was given, this was not noticed after Cephaeline.—*L* '95, II 1276; *P J* '95, I 435.

A further communication (*B M J E*, '05, I 12) on these interesting alkaloids shows that qualitatively the action of Cephaeline is the more intense. As expectorants the alkaloids rank with Senega and Squills.

Dose — $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.008 to 0.0018 gramme, as an expectorant; $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.008 to 0.0018 gramme, as an emetic

Tests — Emetine Hydrochloride dissolves readily in Water. The solution yields the distinctive tests with Potassium or Sodium Hydroxide Solution given under Emetine Hydrochloride. The separated alkaloid should conform to the distinctive tests given under Emetine. The aqueous solution faintly acidified with Nitric Acid yields with Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, but which, when washed, is soluble in Ammonia Solution and Potassium Cyanide Solution. The salt should leave no weighable residue when ignited with free access of air.

VINUM EMETINÆ — This wine should contain $3\frac{1}{2}$ grains of Emetine Hydrochloride in 8.1 oz. or be equal to Vinum Ipecacuanhæ, B.P.

CEPHAELINE — $C_{11}H_{19}NO$, eq 232.44 or $C_{11}H_{19}N_2O$, 464.88, the alkaloid discovered by Paul and Cowley in both Brazilian and Columbian Ipecacuanha.

Colourless, crystalline alkaloid which, like Emetine, is exposure to light. It is readily soluble in alcohol and in alkali less soluble in Ether than Emetine. It forms crystalline salts with acids.

The B.P. Code differs from Paul and Cowley in stating that Cephaeline is more soluble in Ether than Emetine.

Tests — Cephaeline, when crystallised from its concentrated solution in Ether in the presence of Water, melts at 96° to 98° C. (204.8° to 208.4° F.), when crystallised by the addition of Ammonia to a salt in the presence of Ether it melts at 102° C. (215.6° F.). It neutralises acids with the formation of salts which are neutral in reaction towards the ordinary indicators of neutrality. It may therefore be titrated with Normal Volumetric Hydrochloric Acid Solution, using either Methyl Orange, Hematoxylin or Iodocyan Solution as an indicator. 1 c.c. of Normal Volumetric Hydrochloric Acid Solution is equivalent to 0.23244 gramme of Cephaeline. All (xxvii 345) gives the following colour reactions for Cephaeline.

— With Ferric Chloride, Cephaeline from Rio Ipecacuanha gives a bluish-green coloration, the alkaloid from Carthage gives an indefinite reaction, with Starch and Iodic Acid containing 0.5 per cent. of Iodic Acid (Froehde's reagent), Cephaeline from the Rio root gives a pink colour changing to green, that from the Carthage root a reddish-purple colour, with Froehde's reagent and Hydrochloric Acid a Prussian blue colour, with Starch and Iodic Acid a negative reaction, with Ferric Chloride and Potassium Ferricyanide, Cephaeline from Rio root gives an almost immediate blue coloration, whilst that from Carthage root yields an immediate blue.

CEPHAELINE HYDROCHLORIDE — Readily soluble in Water. In the dry state it has the formula $C_{11}H_{19}NO \cdot HCl$, eq 268.63, but when crystallising from a slightly acid solution, it approximates to $C_{11}H_{19}NO \cdot HCl \cdot 3H_2O$, eq 322.27.

Medicinal Properties — Cephaeline is more powerfully emetic than Emetine, and does not produce purgative effects in doses of $\frac{1}{10}$ to $\frac{1}{2}$ grain = 0.008 to 0.01 gramme, but is slow in action. — L'95, ii 1274

Tests — Cephaeline Hydrochloride dissolves readily in Water. The separated alkaloid should yield the distinctive tests given under Cephaeline. The aqueous solution faintly acidified with diluted Nitric Acid yields on the addition of Silver Nitrate Solution a white curdy precipitate, insoluble in Nitric Acid and, when washed, soluble in Ammonia Solution and in Potassium Cyanide Solution. The salt should leave no weighable residue when ignited with free access of air.

PSYCHOTRINE — Pale lemon-yellow coloured, well-defined transparent prisms. Insoluble in Water, readily soluble in Alcohol or in Chloroform, the solutions becoming a yellow colour on exposure to light, and depositing a dark brown substance.

Tests — Psychotrine melts at 138° C. (280.4° F.). It combines with acids to form salts which are neutral in reaction towards the ordinary indicators of neutrality. It may therefore be determined by titration with Normal Volumetric Hydrochloric Acid Solution using either Methyl Orange, or Iodocyan Solution as an indicator. It appears to have a much higher molecular weight than either Emetine or Cephaeline.

According to Allen (*Analyst*, xxvii 349), Pyschotrine gives the following colour reactions —With Ferric Chloride the alkaloid from Rio root gives a pale cherry red coloration, that from Carthage root an indefinite reaction, with Sulphuric Acid containing 0.5 p.c. of Molybdic Acid (Froehde's reagent), Cephaeline from Rio root gives a pale pink coloration, that from Carthage root a dull purple, with Froehde's reagent and Hydrochloric Acid the alkaloid from either variety gives a pale pink changing to pale green, with Starch and Iodic Acid a blue coloration is produced, the colour being more marked in the case of the alkaloid from Rio root than with that from Columbian root, with Ferric Chloride and Potassium Ferricyanide, the Psychotrine from either variety of root gives an immediate blue coloration. The most valuable means of detecting Ipecacuanha alkaloids consists in the production of Cephaeline in a crystalline form. It is readily obtained by shaking out an Amyl Alcohol or Chloroform solution of the alkaloid with a little dilute Acetic Acid. The acid liquids separated, concentrated, if necessary, and placed on a microscope slide furnished with a coil. A watch glass or small beaker moistened with Ammonia Solution is inverted over the alkaloidal acetate solution, when the absorbed Ammonia vapour liberates the alkaloid in characteristic crystals, which can be observed under the microscope.

Not Official

IRIS

The Rhizome and Roots of *Iris versicolor*, L.

Medicinal Properties —The preparation, Iridin and Extractum Iridis are purgative and diuretic. Emetic and cathartic in large doses. Used in biliousness, torpid liver and duodenal dyspepsia.

IRIDIN —A dark brown powder, obtained from Iris.

Dose —1 to 5 grains = 0.065 to 0.32 gramme. Given in pill with Extract of Henbane, but more usually combined also with Eucalyptin and other cholagogues.

'Diluted Glucose' is a good excipient for Iridin.

It has been known for many years as an eclectic remedy, under the names Iridin and Iridin.

Extractum Iridis of pilular consistence prepared with Alcohol (94 p.c.) was Official in *U.S.P.*, 1890, but was omitted in 1900, and a powdered extract prepared with Alcohol (60 p.c.) was included in *B.P.C. Formulary* 1901, with the synonym Iridin, this has been incorporated in the *B.P.C.*

Not Official

ISPAGHULA

The dried Seeds of *Plantago ovata*, Forsk.

Demulcent, and mildly astringent. They are given (whole) in protracted diarrhoea in India. In their passage through the alimentary canal they absorb Water, swell up and yield a bland demulcent mucilage. Official in the *Ind* and *Col Add* for India and the Eastern Colonies.

Dose.—50 to 150 grains = 3.2 to 10 grammes.

DECOCTUM ISPAGHULÆ.—Boil 120 grains of bruised Ispaghula with 24 fl. oz. of Distilled Water for 10 minutes, strain, and make up to 20 fl. oz. by rinsing contents of strainer with more of the Water, if necessary.

Dose.—1 to 2 fl. oz. = 14.2 to 56.8 c.c.

This is official in the *Ind* and *Col Add* for India and the Eastern Colonies. It has been incorporated in the *B.P.C.*

Not Official.

IZAL.

A distilled product from coke, introduced as a non-pyogenic disinfectant, and sold in three forms: (1) medical Izal, (2) an emulsion containing 40 p.c. of the refined oil, (3) pure Izal, an emulsion containing 40 p.c. of unrefined oil.

Izal in the treatment of phthisis 10 minims = 0.6 cc, mixed with Cod-liver Oil given internally, and as an inhalation by evaporation at the bedside, and as a solution in paraffin used as a spray — *L* '02, 1 146

Intra-tracheal injection of Izal Oil in phthisis, 60 minims of a 10 p.c. solution of Izal Oil in Glycerin — *B M J* '02, 1 479, *Trans Brit Cong Tub*, 111 413, also Izal, 20 minims in Glycerin, 1 oz, with occasionally 10 minims Guaiacol added — *B M J* '03, 1 545

Recommended (*B M J* '04, 11 1520) in the treatment of ringworm, the pure Izal being well rubbed into the scalp.

In dysentery — *I M G* '05, 11 261. From 1 to 2 drms in a pint of Water at 100-104° F (37.7 to 40° C). From 1 to 2 pints of the solution are run into the large gut, the injection being retained from 10 to 15 minutes. The treatment should be carried out twice daily in acute cases and once in subacute and chronic cases. Fifteen to 25 minims six or seven times a day in dysentery (*I M G* '05, 11 281), the drug being made up with Chloric Ether, Cardamoms and Glycerin or with Spirits of Chloroform and Peppermint.

Izal Oil as an antiseptic disinfectant, given in doses of 1 to 3 capsules containing 2 minims = 0.12 cc of Izal Oil in each.

JABORANDI FOLIA.

JABORANDI LEAVES

The dried Leaflets of *Pilocarpus Jaborandi*, Holmes

The Jaborandi Leaves of commerce have been very variable, and are the produce of different varieties of *Pilocarpus*.

The principal alkaloid is Pilocarpine, a syrupy liquid, forming crystalline salts, the Hydrochloride and Nitrate are most used, see pp 893, 895. They also contain Isopilocarpine, which is not so powerful a weaker properties, and from 0.2 p.c. to 1.1 p.c. of an alkaloid of weaker properties and Isopilocarpine are isomeric. The *B P* does not require the leaves to conform to any definite alkaloidal standard. They vary considerably in the amount of Pilocarpine they contain, generally about 0.5 p.c. It is still an open question whether in the present state of our knowledge of the alkaloids, the standardisation can be justified. Jowett (*Y B P* '99, 135, *C D* '99, 1 203) is of opinion that it is possible to determine the alkaloid with a fair degree of accuracy. The information is, however, of little value, for it gives no indication of the amount of Pilocarpine contained in the total alkaloids, and it must be assumed that on the Pilocarpine alone depends the therapeutic value of the preparation.

It has been recommended that, if the drug is retained, *Pilocarpus microphyllus* should be substituted for the present official variety, and that the galenic preparations be standardised. The limit suggested for the leaves is 0.5 to 0.75 p.c. of total alkaloid. The ratio of Pilocarpine to other alkaloids appears to be practically constant in this variety.

The Leaflets of *Pilocarpus Jaborandi*, Holmes, and of *Pilocarpus microphyllus*, Stapf, are official in the *U S P*, and are required to yield not less than 0.5 p.c. of alkaloids.

Jaborine is a mixture of Pilocarpine, Isopilocarpine, and, possibly, a trace of Pilocarpidine, with a trace of colouring matter.

Medicinal Properties—Powerful and prompt diaphoretic, sialagogue, and galactagogue. Useful in the dropsy, uræmia and thirst of Bright's disease. It is antagonistic in its action to Belladonna. The salts of Pilocarpine, owing to their more constant action, are more generally used than the galenical preparations of Jaborandi. See also Pilocarpine Nitras.

Official Preparations—Extractum Jaborandi Liquidum and Tinctura Jaborandi. Used in the preparation of Pilocarpine Nitras.

Foreign Pharmacopœias—Official in Austr., Belg., Fr., Ger., Ital., Jap., Mex., Port., Span. and Swiss, U.S. (Pilocarpus). Not in the others.

Descriptive Notes.—The Leaves of *Pilocarpus Jaborandi*, Holmes, official in the *B.P.*, are no longer obtainable in commerce. They are described as dull green, oval oblong or oblong lanceolate, from 2½ to 4 in (6 to 10 cm.) in length (8 to 16, mostly 12, mm., *P.G.*), 12 cm. long and 3 to 4 in broad, *U.S.P.*, obtuse and emarginate at the apex, and unequal and shortly petiolate at the base, with an entire, slightly revolute margin, of a coriaceous texture, glabrous, or with a few scattered hairs on the under surface, and with the lateral veinlets distinctly prominent on the upper surface, containing numerous oil glands, and having an aromatic odour, pungent taste, and increasing the flow of saliva when chewed.

The Leaves which are now in commerce are those of *Pilocarpus pennatifolius*, Lam., from Paraguay, which are very similar in size and shape, but have a greyish-green colour, the lateral veinlets are scarcely prominent on the upper surface, and the base of the leaf is usually equal and tapering. The leaves of *P. trachylophus*, Holmes, are similar in shape to those of *P. Jaborandi*, but rather smaller, with prominent veins on the upper surface and revolute margins, but are of a brownish green tint, and covered on the under surface with curved simple hairs, and there are usually only two pairs of leaflets on the leaves besides the terminal one, whereas in *P. Jaborandi* and *P. pennatifolius* there are three or four pairs. The leaflets of *Pilocarpus microphyllus*, Staff., are very much smaller (1.2 to 3.7 cm. long, 0.8 to 1.6 cm. broad, *U.S.P.*), the lateral ones sessile, rhomboid oval, or obovate obtuse and emarginate, with pellucid glands, veins coarsely reticulated, but not very prominent, almost odourless, but resemble Jaborandi in taste. These leaves contain about the same quantity of Pilocarpine as those of *P. Jaborandi*, and are largely used for the preparation of the alkaloid, they are the only kind that can replace the official leaves of the *B.P.* so as to give a preparation of equal strength. Unfortunately, there is a spurious leaf very like it offered in the market at intervals, derived from *Swartzia decumbens*, Holmes, a leguminous plant, not possessing the properties of Jaborandi. These may be distinguished by having very short hairy stalks about 1 mm. long, by the veinlets being more or less translucent, and by the presence of smaller rounded leaflets mixed with the ovate or oval leaflets. See *P.J.* (4) in 2. Other varieties of Jaborandi which have appeared in the market, but not regularly, are described in *P.J.* (4) 1: 501,

Not Official

IZAL

A distilled product from coke, introduced as a non-poisonous disinfectant, and sold in three forms (1) medical Izal, (2) an emulsion containing 40 p c of the refined oil, (3) ordinary Izal, an emulsion containing 40 p c of unrefined oil.

Izal in the treatment of phthisis 10 minims = 0.6 c c, mixed with Cod-liver Oil given internally, and as an inhalation by evaporation at the bedside, and as a solution in paraffine used as a spray — *L* '02, i 146

Intra-acheal injection of Izal Oil in phthisis, 60 minims of a 10 p c solution of Izal Oil in Glycerin — *B M J* '02, i 479, *Trans Brit Cong Tub*, iii 413, also Izal, 20 minims in Glycerin, 1 oz, with occasionally 10 minims Guaiacol added — *B M J* '03, i 545

Recommended (*B M J* '04, ii 1520) in the treatment of ringworm, the pure Izal being well rubbed into the scalp

In dysentery — *I M G* '05, ii 261. From 1 to 2 drms in a pint of Water at 100-104° F (37.7 to 40° C) From 1 to 2 pints of the solution are run into the large gut, the injection being retained from 10 to 15 minutes. The treatment should be carried out twice daily in acute cases and once in subacute and chronic cases. Fifteen to 25 minims six or seven times a day in dysentery (*I M G* '05, ii 281), the drug being made up with Chloric Ether, Cardamoms and Glycerin or with Spirits of Chloroform and Peppermint

Izal Oil as an intestinal disinfectant, given in doses of 1 to 3 capsules containing 2 minims = 0.12 c c of Izal Oil in each

JABORANDI FOLIA.

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The dried Leaflets of *Pilocarpus Jaborandi*, Holmes

The Jaborandi Leaves of commerce have been very variable, and are the produce of different varieties of *Pilocarpus*

The principal alkaloid is Pilocarpine, a syrupy liquid, forming crystalline salt- the Hydrochloride and Nitrate are most used, *see* pp 893, 895. They also contain which possesses similar but weaker properties, an ethereal oil. Pilocarpine and its derivatives are isomeric. The *B P* does not require the leaves to conform to any definite alkaloidal standard. They vary considerably in the amount of alkaloid they contain, generally about 0.5 p c. It is still an open question as to in the present state of our knowledge of the alkaloids, the standardisation can be justified. Jowett (*Y B P* '99, 435, *C D* '99, i 203) is of opinion that it is possible to determine the amount of alkaloid with a fair degree of accuracy. The information is, however, of little value, for it gives no indication of the amount of Pilocarpine contained in the total alkaloids, and it must be assumed that on the Pilocarpine alone depends the therapeutic value of the preparation.

It has been recommended that, if the drug is retained, *Pilocarpus nurophyllus* should be substituted for the present official variety, and that the galenic preparation be standardised. The limit suggested for the leaves is 0.5 to 0.75 p c of total alkaloid. The ratio of Pilocarpine to other alkaloids appears to be practically constant in the varieties.

The Leaflets of *Pilocarpus Jaborandi*, Holmes, and of *Pilocarpus nurophyllus*, Stapf, are official in the *U S P*, and are required to yield not less than 0.1 p c of alkaloids.

Jaborine is a mixture of Pilocarpine, Isopilocarpine, and, possibly, a trace of Pilocarpidine, with a trace of colouring matter.

Medicinal Properties—Powerful and prompt diaphoretic, sialagogue, and galactagogue. Useful in the dropsy, uræmia and thirst of Bright's disease. It is antagonistic in its action to Belladonna. The salts of Pilocarpine, owing to their more constant action, are more generally used than the galenical preparations of *Jaborandi*. See also *Pilocarpine Nitras*.

Official Preparations *Extractum Jaborandi*, *Liquidum* and *Tinctura Jaborandi*. Used in the preparation of *Pilocarpine Nitras*.

Foreign Pharmacopœias—Official in Aust., Belg., Fr., Ger., Ital., Jap., Mex., Port., Span. and Swiss, U.S. (*Pilocarpus*). Not in the others.

Descriptive Notes The leaves of *Pilocarpus Jaborandi*, Holmes, official in the *B.P.*, are no longer obtainable in commerce. They are described as dull green, oval oblong or oblong lanceolate, from 2½ to 4 in (6 to 10 cm) in length (8 to 16, mostly 12, cm, *P.G.*), 12 cm long and 3 to 4 in broad, *U.S.P.*), obtuse and emarginate at the apex, and unequal and shortly petiolate at the base, with an entire, slightly revolute margin, of a coriaceous texture, glabrous, or with a few scattered hairs on the under surface, and with the lateral veinlets distinctly prominent on the upper surface, containing numerous oil glands, and having an aromatic odour, pungent taste, and increasing the flow of saliva when chewed.

The leaves which are now in commerce are those of *Pilocarpus pennatifolius*, Hemsl., from Paraguay, which are very similar in size and shape, but have a greyish-green colour, the lateral veinlets are scarcely prominent on the upper surface, and the base of the leaf is usually equal and tapering. The leaves of *P. trachylophus*, Holmes, are similar in shape to those of *P. Jaborandi*, but rather smaller, with prominent veins on the upper surface and revolute margins, but are of a brownish-green tint, and covered on the under surface with curved simple hairs, and there are usually only two pairs of leaflets on the leaves besides the terminal one, whereas in *P. Jaborandi* and *P. pennatifolius* there are three or four pairs. The leaflets of *Pilocarpus macrophyllus*, Staff., are very much smaller (1.2 to 3.7 cm long, 0.8 to 1.6 cm broad, *U.S.P.*), the lateral ones sessile, rhomboid oval, or obovate obtuse and emarginate, with pellucid glands, veins coarsely reticulated, but not very prominent, almost odourless, but resemble *Jaborandi* in taste. These leaves contain about the same quantity of Pilocarpine as those of *P. Jaborandi*, and are largely used for the preparation of the alkaloid, they are the only kind that can replace the official leaves of the *B.P.* so as to give a preparation of equal strength. Unfortunately, there is a spurious leaf very like it offered in the market at intervals, derived from *Swartzia decipiens*, Holmes, a leguminous plant, not possessing the properties of *Jaborandi*. These may be distinguished by having very short hairy stalks about 1 mm long, by the veinlets being more or less translucent, and by the presence of smaller rounded leaflets mixed with the ovate or oval leaflets. See *P.J.* (4) iii 2. Other varieties of *Jaborandi* which have appeared in the market, but not regularly, are described in *P.J.* (4) i 501,

(4) xvii 713 The last, which comes from Guadalupe, contains as much Pilocarpine as the Paraguay Jaborandi. It is derived from *Pilocarpus racemosus*, Vahl. It has larger, broader, and somewhat obovate leaves. The *PG* gives as a distinguishing feature of Jaborandi Leaves that the palisade cells should be about one-fifth of the thickness of the leaf. In powder, Jaborandi may be recognised by polygonal epidermal cells with a strongly striated cuticle, thick-walled bast fibres, one-celled hairs, seriate cluster crystals and the palisade cells. The epidermal cells of *Swartzia decipiens* are very sinuous, and there are pluri-cellular hairs, the terminal cells being largest, on the nerves. Recently the leaves of a species of *Casearia*, Nat. Ord. Sauriadea, have been offered as Jaborandi. They are oblong, elliptic, tapering towards both ends, thinner, and have linear, as well as round, oil receptacles in the leaves.

Tests—The method adopted by the *USP* for the assay of the Leaves is essentially as follows—A weighed quantity of 10 grammes of the Leaves in No 60 powder is moistened with 2 c c of Ammonia Solution and 3 c c of Chloroform, and packed firmly in a small cylindrical percolator provided with a pledget of cotton-wool firmly packed in the neck, and slowly percolated with Chloroform containing about 2 p c of Ammonia Solution, until exhausted, about 100 c c usually being sufficient. The percolate is transferred to a separator, and the alkaloids removed by shaking with 15 c c of Normal Volumetric Sulphuric Acid Solution, the acid liquid being separated and transferred to a second separator, the extraction of the alkaloidal residue being continued with a second quantity of a mixture of 2 c c of Normal Volumetric Sulphuric Acid Solution and 8 c c of Water, followed by 10 c c of Water, the aqueous acid portion and the aqueous portion being separated in each instance and transferred to the second separator. After the addition of sufficient Ammonia Solution to render the liquid alkaline, the liberated alkaloids are extracted by shaking with three successive quantities of 20 c c, 15 c c and 10 c c of Chloroform, the chloroformic solution separated in each instance, transferred to a beaker or flask, the Chloroform evaporated at a gentle heat, the alkaloidal residue dissolved in 7 c c. of Tenth-normal Volumetric Sulphuric Acid Solution and the excess titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, using Cochineal or Iodeosin Solution as an indicator of neutrality. The number of c c of Fiftieth-normal Volumetric Potassium Hydroxide Solution used divided by 5, the quotient subtracted from 7, the remainder multiplied first by 0.02 and then by 10, yields the percentage of total alkaloids in terms of Pilocarpine present in the sample. The Leaves yield from 1 to 7 p c of ash.

Preparations

EXTRACTUM JABORANDI LIQUIDUM LIQUID EXTRACT OF JABORANDI

20 of Jaborandi Leaves, in No 20 powder, percolated with Alcohol (45 p c), until 67 volumes have been obtained. Reserve the

first 17 and evaporate the remainder to a soft extract, which is dissolved in the first portion and made up with Alcohol (45 p c) to 20 (1 in 1)

The Liquid Extract of Jaborandi official in the *B P* is not a standardised preparation, though it has been recommended that, if retained, it should be standardised and a method of assay given. The *U S P* Fluid Extract is required to contain 0.4 p c w/v of the alkaloids from *Pilocarpus*. The *P G* does not include a Fluid Extract.

Dose 5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—official in *U S* Fluidextractum *Pilocarpi*, 1 in 1 and standardised, *B P* '85 had a solid **Extractum Jaborandi** prepared with Alcohol (57 p c), and *L P C* have adopted this, using Alcohol (60 p c), *B P C* have also **Infusum Jaborandi**, 1 in 20 of boiling Water, infused 15 minutes. Not in the others.

Tests—Liquid Extract of Jaborandi has a sp gr of 1.010 to 1.040, contains from 12.0 to 22.0 p c w/v of total solids, and about 34 p c w/v of Absolute Alcohol. As mentioned above, the official preparation is not standardised, and no method is given for the determination of the total alkaloids. It may be conveniently assayed by the method suggested by Farn and Wright (*Y B P* '99, 381, *C D* '99, n 205). A measured quantity of 10 c c is acidified with dilute Sulphuric Acid and evaporated on a water bath to a syrupy consistence, 30 c c of Alcohol (90 p c) added, and the mixture well stirred and allowed to stand for an hour. The liquid portion is then separated by decantation or filtration, the mucilaginous deposit dissolved in a little acidulated Water, and the treatment with Alcohol repeated. The dish and filter are rinsed with a little Alcohol, the filtrates and rinsings bulked and evaporated over a water-bath, Water being added from time to time until all the Alcohol has been removed. The residual liquor is transferred to a separator, the dish washed with a few drops of Water, and the whole rendered alkaline with Ammonia Solution. The liberated alkaloids are shaken out with two successive quantities, each of 15 c c, of Chloroform, the chloroformic solutions are separated in each case, mixed, and the alkaloids extracted by shaking with three successive quantities of 9 c c each of a mixture of 25 c c of Water and 2 c c of Semi-normal Volumetric Sulphuric Acid Solution. The acid solutions are separated in each case, mixed, rendered alkaline with Ammonia Solution, and the liberated alkaloids shaken out with two successive quantities each of 15 c c of Chloroform. The chloroformic layer is separated in each case, the liquids mixed, the Chloroform evaporated over a water-bath, the alkaloids dried and weighed. The residue is dissolved in a little Alcohol (90 p c), a calculated excess of Tenth normal Volumetric Hydrochloric Acid Solution and some Water added, and the excess of Volumetric Acid titrated with Twentieth normal Volumetric Sodium Hydroxide Solution, using Cochineal Solution as an indicator of neutrality. Specimens of the Liquid Extract examined in the author's laboratory by the above process gave gravimetrically 0.2, 0.39, and 0.26 p c w/v of total alkaloids, the volumetric determination giving 0.14, 0.3, and 0.16 p c w/v. In performing the volumetric determination, a small

deviation was made from Farr and Wright's method. The alkaloidal residue was dissolved in a measured excess of Tenth-normal Volumetric Hydrochloric Acid Solution, and the excess of Volumetric Acid determined by titration with Hundredth-normal Volumetric Sodium Hydroxide Solution, using Iodeosin Solution as an indicator of neutrality. Farr and Wright, in the examination of 12 specimens of the Liquid Extracts, found from 0.03 p.c. w/v to 0.24 p.c. w/v, and an average of 0.15 p.c. w/v of total alkaloids.

The method of determination adopted by the *USP* is essentially as follows:—A measured quantity of 10 c.c. is dropped on to a little clean sand, contained in a porcelain evaporating basin, and evaporated to dryness on a water-bath. The extract is mixed uniformly with the sand, transferred to an Erlenmeyer flask, the dish rinsed with a mixture of 25 c.c. of Chloroform and 2½ c.c. of Ammonia Solution, which is transferred to the flask, and the whole well shaken at intervals for an hour. The liquid is decanted into a separator, the residue washed with several portions of Chloroform, which are drawn off and filtered into the separator. The alkaloids are extracted from the chloroformic solution by shaking first with 15 c.c. of Normal Volumetric Sulphuric Acid Solution, then with a mixture of 5 c.c. of a similar Volumetric Acid solution and 5 c.c. of Water, and finally with 10 c.c. of Water. The acid, acid and aqueous, and aqueous shakings are separated, transferred to a second separator, rendered alkaline with Ammonia Solution, and the liberated alkaloids shaken out with three successive quantities of 20 c.c., 15 c.c., and 10 c.c. each of Chloroform. The chloroformic layer is separated in each case, the chloroformic solutions mixed, the Chloroform evaporated on a water-bath, and the alkaloidal residue is dissolved in 8 c.c. of Tenth-normal Volumetric Sulphuric Acid, the excess of Volumetric Acid being titrated with 1 c.c. of Volumetric Potassium Hydroxide Solution. The number of c.c. required divided by 5, the quotient subtracted from 8 and the remainder multiplied first by 0.02 and then by 10, yields the percentage w/v of total alkaloids in terms of Pilocarpine present in the Fluid Extract. It may be noticed in this instance that an error appears in the *USP*, 5 c.c. was mentioned as the quantity of Tenth-normal Volumetric Sulphuric Acid Solution to be used in dissolving the alkaloidal residue, whereas in the subsequent calculation of the volumetric result, the quotient obtained by dividing the number of c.c. of excess Fifthieth-normal Volumetric Potassium Hydroxide Solution 5, was directed to be subtracted from 8. This was corrected in the Additions and Corrections (1907).

In criticising a paper on standards of medicines, Jowett (*P. J.* '02, ii 672) states that he, in conjunction with Professor Marshall, has shown, he trusts conclusively, that the galenical preparations of Jaborandi were unreliable, and, furthermore, unnecessary, as the whole therapeutic activity of Jaborandi Leaves can be obtained by Pilocarpine, which might, therefore, be used in place of any galenical preparation. They show the fallacy of attempting to standardise the Liquid Extract on a percentage of total alkaloids. Of two specimens of Liquid Extract yielding 0.21 p.c. and 0.25 p.c. of total alkaloid respectively, one

yielded no crystalline Pilocarpine Nitrate and the other 0.082 per cent. They claim that Pilocarpine Nitrate should be the official salt and all galenical preparations of Jaborandi abandoned.

Farr and Wright (*P. J.* '03, 19) do not admit Jowett's conclusions, and consider that they are not justified by the facts recorded in the paper referred to in his letter. The titration figures of the alkaloid residues obtained in their research based on the molecular weight of Pilocarpine, were in close accord with the gravimetric results showing that the alkaloid was practically pure. They consider there is absolutely nothing in Jowett and Marshall's work to show that any one of the four samples of Liquid Extract used for the physiological experiments was the official article. Judging from the fact that two of them apparently contained little or no Pilocarpine, and the other two quite trifling proportions, it is tolerably certain that they were commercial products of a spurious Jaborandi, and not of the official variety. So long as the demand for galenicals exists, so long will standardisation be required. For the great majority of preparations of alkaloidal drugs the best, because the most natural standard, is one of total alkaloids. Jowett in reply (*P. J.* '03, 141) shows that the case for the expulsion of galenical preparations of Jaborandi does not rest only on two assumptions but points out that the physiological action of Jaborandi is fully produced by pure Pilocarpine, the purity being assured by constancy of the mp and specific rotation after repeated crystallisation. Farr and Wright's statement that the titration figures based on the molecular weight of Pilocarpine were in close accord with the gravimetric results, showing that the alkaloid was practically pure, indicate that they ignore the existence of Isopilocarpine as well as that of Pilocarpidine. Isopilocarpine, although isomeric with Pilocarpine, possesses but one-eighth of the physiological activity of the latter, and would give precisely the same figures on titration. The alkaloid which Farr and Wright assume to have been practically pure might, therefore, from their results, have consisted entirely of Isopilocarpine and contained no Pilocarpine. With regard to the Liquid Extracts, the four preparations were guaranteed as such by three prominent wholesale druggists, and such as would be largely distributed to the pharmacists of this country. The object was to determine the therapeutic value of the preparations of Jaborandi such as would be used in dispensing, for which purpose the specimens referred to were purchased from three manufacturing houses of the highest repute.

Farr and Wright (*P. J.* '03, 171) still maintain that, as isolated by the process employed by them, the total alkaloids consist of almost pure Pilocarpine. They consider that the proportion of alkaloid, other than Pilocarpine is so small that it may be safely ignored, and quote the results of Jowett's examination of the genuine drug. With regard to the galenical preparations, they think that the explanations given in his letter have not improved his position. It is certain that the official Jaborandi is an exceedingly active drug, and equally certain that the same degree of activity will be manifested by a Liquid Extract if carefully prepared. They repeat that the research

does not justify the wholesale denunciation of the galenical preparations of the official drug

TINCTURA JABORANDI. TINCTURE OF JABORANDI

4 Jaborandi Leaves, in No 40 powder, percolated with Alcohol (45 p c), to yield 20

The Official Tincture is not a standardised preparation The *USP* does not include a Tincture

Dose —30 to 60 minims = 1·8 to 3 6 c c

Foreign Pharmacopœias Official in Fr and Mex, 1 in 5, Span, 1 in 10 Not 1

Wright and Farr (*PJ* (3) xxii 1) show an enormous variation in the strength of various samples of this tincture, viz, from 0 032 to 0 148 p c of alkaloid, and recommend a standard of 0 1 p c

Tests.—Tincture of Jaborandi has a sp gr of 0 950 to 0 960, contains from 2 5 to 3 5 p c w/v of total solids, and about 40 p.c. w/v of Absolute Alcohol A standard of 0 048 p c w/v of total alkaloids has been suggested for the Tincture

A convenient method for the determination of the total alkaloids is that of Farr and Wright With the exception that a measured quantity of 50 c c of the Tincture is substituted for 10 c c of the Liquid Extract, the process may be conducted as described under the heading of Extractum Jaborandi Liquidum

A Tincture prepared in the author's laboratory had a specific gravity of 0 955, contained 2·5 p c w/v of total solids and 40 3 p c w/v of Absolute Alcohol When assayed according to the method mentioned above, it yielded gravimetrically 0 036 p c w/v of alkaloids, which on titration showed 0 036 p c w/v reckoned as Pilocarpine

PILOCARPINÆ NITRÆ. See p 893

JALAPA.

JALAP

Fr, JALAP, GER, JALAPENWURZEL, ITAL, GIALAPPA, SPAN, JALAPA

The dried Tubercules of *Ipomœa Purga*, Hayne

It contains, as its principal ingredient, **Convolvulin**, soluble in Alcohol, but insoluble in Ether, and but a small part of Resin Jalapæ, *BP*

The *BP* requires that Jalap should yield not less than 9·0 p c, nor more than 11 0 p c of Resin answering the official requirements, the *USP* not less than 7 p c of total Resin, of which not more than 15 p c should be soluble in Ether, the *PG* at least 9 0 p c. of Jalap Resin

Medicinal Properties —A brisk cathartic, operating sometimes painfully, producing copious watery discharges from its hydragogue powers, it is especially serviceable in dropsy and cerebral

congestion, when it is usually prescribed in the form of the Compound Powder

Dose—5 to 20 grains=0.32 to 1.3 grammes

Swiss, minimum single dose, 1.0 gramme, maximum daily dose, 5.0 grammes

Prescribing Notes—*The powder can be given in cachets, or mixed with Confections. The Resin is given in pills made by adding 'Diluted Glucose,' q s*

Official Preparations—*Extractum Jalapæ, Pulvis Jalapæ Compositus, Jalapæ Resina, Tinctura Jalapæ*, used in the preparation of *Pulvis Scammonii Compositus*. The resin is contained in *Pilula Scammonii Composita*

Not Official—*Mistura Jalapæ cum Rheo, Pilula Jalapæ, Tinctura Jalapæ Composita, Sapo Jalapinus, Jalapin*

Foreign Pharmacopœias—Official in Austri, Hung and Swiss, at least 10 p.c. of Resin, Belg., Dutch and U.S., 9 p.c., Dan., Fr., Norw. and Swed., 7 p.c. Ger., Jap. and Russ., 9 p.c., Ital. (Malappa), 12 p.c., Mex., 11 p.c., Span., 15 to 18 p.c.

The Br. Codex (1834) fixed the standard at 15 to 18 p.c. of Resin, lowered in 1908 to 7 p.c., U.S. (1880 and 1900) at 12 p.c., Ger. (1890) lowered the figure to 7 p.c., but (1900) increased it again to 'at least 9 p.c.'

Descriptive Notes—The Jalap of commerce is usually imported from Vera Cruz and consists of ovoid, or more or less broadly fusiform or subspherical roots, averaging about $1\frac{1}{2}$ to 3 in (1 to 3 in, $2\frac{1}{2}$ to $7\frac{1}{2}$ cm B.P.), but is sometimes 4 to 5 in or more in diameter. The larger roots are often incised to facilitate drying. Externally the roots are of a dark greyish-brown colour, furrowed and wrinkled, and marked with numerous short transverse paler scars or lenticels. A transverse section exhibits a yellowish grey or brown tint with irregular darker concentric rings, consisting of Resin cells, it has a smoky odour, and at first a sweetish, then an acrid taste and a disagreeable flavour. There is considerable difference in the density of the roots as met with in commerce, the light pieces containing most Resin, the heavier pieces apparently owing their weight to sugar, which is difficult to entirely separate from the Resin. Jalap has been cultivated in India and Jamaica, and these roots differ from the Mexican in their paler and more starchy appearance internally. The Indian, which shows a tendency to a fusiform shape, is sometimes unusually rich in Resin, the Jamaica Jalap more frequently presents a subglobular form, it has sometimes been imported in the form of transverse slices, but since the comparative disuse of the drug of late years and the consequent fall in price, the exportation from thence has apparently ceased. Powdered Jalap is characterised by the starch grains, often compound and sometimes amorphous from the action of heat, by the laticiferous cells and globules of resin escaped from them, the pitted vessels as well as tracheids, sphaerocrystals of Calcium Oxalate often 2 to 5 in a parenchymatous cell, and sclerenchymatous cells. A variety of Jalap known in commerce as Tampico Jalap, derived from *Ipomœa simulans*, Hanbury, is occasionally imported. It is more fusiform, smaller, more shrunken, and does not exhibit pale transverse lenticels. The root of another species, *Ipomœa Orizabensis*, Ledan, has been recently imported in large quantities under the name of Mexican Scammony Root. This

root is spindle-shaped and about 2 feet long, and occurs in commerce under the name of Stalk Jalap, in regularly rectangular pieces 1 or 2 inches in diameter and 2 to 3 inches long, which exhibit a radiate transverse section, and numerous thick bundles of vessels projecting as stiff fibres from the fractured surface.

Tests—The Resin of Jalap is soluble in Alcohol (90 p c), and this solvent is officially employed for its determination. A weighed quantity of, say, 10 grammes of the Jalap in fine powder is digested at a gentle heat for 24 hours with twice its weight of Alcohol (90 p c), transferred to a percolator, and percolated with Alcohol (90 p c) until nothing further is dissolved. The alcoholic solution is precipitated by the addition of Water, the Alcohol distilled, the residue is transferred whilst hot to a dish, cooled, and the supernatant liquid removed, the Resin washed with hot Water, dried and weighed. It should weigh not less than 0.9 nor more than 1.1 gramme. This weight multiplied by 10 yields the p c w/w of Resin present in the sample.

The *U S P* distinguishes between the Ether-soluble and the Ether-insoluble Resin. A weighed quantity of 10 grammes of Jalap in No. 60 powder is percolated in a well-covered percolator, with Ether [sp gr 0.716 at 25° C (77° F)] until 50 c c of percolate have been obtained. The percolate is transferred to a tared beaker, the Ether evaporated on a water-bath, and the residue weighed. The weight multiplied by 10 gives the p c of Ether-soluble Resin. The percolation is continued until 100 c c of Ether-soluble Resin (94.9 p c) until 100 c c of percolate have been obtained. A measured quantity of 20 c c of this percolate is transferred to a separator, mixed with 20 c c of Chloroform, and shaken for 1 minute with 20 c c of Water. The Chloroform layer is separated, transferred to a tared beaker, the separator washed with 5 c c of Chloroform, and the mixed chloroformic liquids are evaporated to dryness on a water-bath, the residue dried till constant in weight, and weighed. This weight multiplied by 50 gives the percentage of Resin insoluble in Ether. The sum of the two weights represents the total Resin. The *P G* exhausts 1 part by weight of coarsely-powdered Jalap Root for 24 hours at a temperature of 35° to 40° C (95° to 115° F), first with 4 parts by weight of the Alcohol (90 p c) and then with a further 2 parts by weight of the Alcohol. The Alcohol is distilled off, the residue of Resin is washed with warm Water, until it no longer colours the latter. The Resin is dried in the water-oven and weighed. The *French Codex* 1908 adopts a standard of 7 p c of Jalap Resin, which is lower than the *B P* or *P G*, the *U S P* standard of not less than 8 p c of total resin, of which not more than 1.5 p c should be soluble in Ether, was altered by the Additions and Corrections 1907 to not less than 7 p c of total resin, of which not more than 1.5 p c should be soluble in Ether.

The ash of Jalap varies from 4 to 6 p c and should not exceed the latter figure.

The retention of the Resin has been recommended as the present official standard for the percentage of Resin.

Preparations

EXTRACTUM JALAPÆ EXTRACT OF JALAP

Jalap, in coarse powder, 1, Alcohol (90 p c), 5, Distilled Water, 10. A solid Extract prepared by treating the Jalap first with the Alcohol and subsequently with the Water, and combining the two residues into one Extract.

100 lb of Jalap yielded 30 lb of Extract

Dose — 2 to 8 grains = 0.13 to 0.32 gramme

PULVIS JALAPÆ COMPOSITUS COMPOUND POWDER OF JALAP

Jalap, 5, Acid Potassium Tartrate, 9. Ginger, 1. (1 in 3)

Dose — 20 to 60 grains = 1.3 to 4 grammes

Foreign Pharmacopœias — Official in Russ. Jalap 1, Potassium Bitartrate 2, U.S., Jalap 3, Potassium Bitartrate 6, Mex. Not in the others.

JALAPÆ RESINA JALAP RESIN

Extracted from Jalap by exhausting with Alcohol (90 p c), and purified by washing with Water.

Dose — 2 to 5 grains = 0.13 to 0.32 gramme

Ital., maximum single dose, 0.3 gramme; maximum daily dose, 1.0 gramme

Foreign Pharmacopœias — Official in Austri., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Tests Jalap Resin, when powdered, is officially required to yield little or nothing to warm Water and not more than 10 p c to Ether. It is readily soluble in Alcohol (90 p c), insoluble in Turpentine Oil. The *U.S.P.* requires that not more than 15 p c should be soluble in Ether, that it should be soluble in Alcohol (94.9 p c) in all proportions, and that the alcoholic solution should be only faintly acid to blue Litmus paper. It also requires that not more than 35 p c should be soluble in Chloroform. The *B.P.* mentions not more than 10 p c as the Ether-solubility limit. The *P.G.* says readily soluble in Alcohol (90 p c), and not more than 10 p c should be soluble in Chloroform, but makes no reference to Ether solubility. The *U.S.P.* requires that it shall possess an Acid value of not more than 13.93 and a Saponification value of at least 139.35. The *U.S.P.* states that the Resin should not suffer material loss of weight when heated at 100° C (212° F), and that the anhydrous Resin melts about 150° C (302° F). Both the *U.S.P.* and *P.G.* require that the Resin should be completely soluble in 5 times its weight of Ammonia Solution, and when this solution is acidified only a slight turbidity at the most should be produced. The *P.G.* warms the Resin with the Ammonia Solution, and requires, in addition, that the solution shall not gelatinise on cooling. It uses Acetic Acid for acidification, whilst the *U.S.P.* employs Hydrochloric Acid. That the Resin should be soluble in 5 times its weight of Ammonia Water is a reasonable requirement, but few samples will be found to respond to the latter half of the test.

The more generally occurring impurities are Scammony Resin and the Resin of Tampico Jalap, Guaiacum Resin, Colophony, Water,

and soluble impurities Scammony Resin and the Resin of Tampico Jalap are detected by the Ether-solubility test, Guaiacum Resin by Ferric Chloride TS failing to produce any greenish-blue coloration in the Alcohol (90 p c) solution, Colophony greatly increases the Acid value and is also readily detected by the Ammonia test. Water is detected by loss of weight at 100° C (212° F), and soluble impurities by evaporating a filtered aqueous trituration of the Resin to dryness, the filtered liquid should be colourless, and no solid residue should remain.

Water—Jalap Resin when triturated with 10 parts of Water should give an almost colourless filtrate, *P G*, Water should not become coloured by it nor dissolve any portion of it, *U S P*

Chloroform—Not more than 35 p c of the Resin should be soluble in Chloroform, *U S P*, if 1 gramme be warmed with 10 grammes of Chloroform and the product filtered, the filtrate after evaporation should not leave a residue of more than 0.1 gramme, *P G*

Ammonia—If Jalap Resin be warmed in a well-closed vessel with 5 parts of Ammonia Solution, a solution should be obtained which, on cooling, is not gelatinous, and on evaporation leaves a residue soluble in Water, all but an insignificant resinous portion. On supersaturating the solution with dilute Acetic Acid a faint turbidity at most should be produced, *P G*, slowly but completely soluble in 5 parts (by weight) of Ammonia Water, and when this solution is acidified with Hydrochloric Acid only a slight turbidity should appear, *U S P*

Ferric Chloride—A few drops of Ferric Chloride TS added to some of the powdered Resin, moistened with Alcohol, should produce no greenish-blue colour, *U S P*

Acid Value—1 gramme of Jalap Resin dissolved in 50 c c of Alcohol containing 1 c c of Phenolphthalein TS should require not more than 0.5 c c of Semi-normal Alcoholic Potassium Hydroxide Volumetric Solution to produce a red colour (limit of acid resins), *U S P*

Saponification Value—If to 1 gramme of Jalap Resin dissolved in 50 c c of Alcohol in a flask, 25 c c of Semi-normal Alcoholic Potassium Hydroxide Volumetric Solution be added and the mixture be heated on a water-bath for one hour, and if the excess of Alkali be titrated with Semi-normal Volumetric Sulphuric Acid Solution, using 5 drops of Phenolphthalein TS as indicator, at least 20 c c of Semi-normal Sulphuric Acid Volumetric Solution should be required, *U S P*

TINCTURA JALAPÆ, TINCTURE OF JALAP

A Tincture obtained by treating Jalap with Alcohol (90 p c), and standardising it to contain 1.5 of the Resin in 100 cc, which is equal to 1 of Root in 6 or 7 of Alcohol (90 p c).

Dose.— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Foreign Pharmacopœias—Official in Belg, 1 in 50 from Resin, Port, 1 and 5 by weight. Not in the others.

Tests.—Tincture of Jalap has a sp gr of 0.905 to 0.910, contains about 3.5 p c w/v of total solids, and about 68.0 p c w/v of Absolute Alcohol. It contains about 1.5 p c of Jalap Resin.

Not Official

MISTURA JALAPÆ CUM RHEO—Jalap Resin, $\frac{1}{2}$ grain, Compound Tincture of Rhubarb, 10 minims, Tragacanth, $\frac{1}{2}$ grain, Syrup of Ginger, 5 minims, Glycerin, 10 minims, Caraway Water, to 1 fl drm. Powder the Resin, mix with the Tragacanth, add the Tincture and then the

other ingredients in the order given Dose —1 fl drm for a child 1 year old —
St Thomas's

Note —The official extract of Jalap varies considerably in strength, hence the Resin of Jalap is used, with Tragacanth to suspend it

This has been incorporated in the *B P C*

TINCTURA JALAPÆ COMPOSITA —Jalap, 8, Scammony, 2, Turpeth,* 1, Alcohol (60 p c), to 100

Dose — $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Official in the *Ind* and *Col Add* for India, the Eastern Colonies, and North American Colonies

This has been incorporated in the *B P C*

Fr, Mex, Port and Span, similar to above, Belg, Jalap 1, Scammony 1.5, Tincture of Ginger, 2.5, Alcohol (80 p c) 950, Swiss, Jalap 10, Scammony 10, Diluted Spirit, to 100

PILULA JALAPÆ —Jalap Soap, 3, Powdered Jalap, 1 —*Ger*

SAPO JALAPINUS —Resin of Jalap, 1, Soap, 1 —*Ger, Jap* and *Swiss*

JALAPIN A purified Resin of Jalap, entirely soluble in Ether

Dose 1 to 5 grains = 0.06 to 0.32 gramme

Tampico Jalap from *Ipomœa similans*, Harb., and Orizaba root (Woody Jalap), from *Ipomœa Orizabensis*, Ledan, also yield a glucoside Jalapin, soluble in Ether, and almost, if not completely, identical with Resin Scammony, *B P*, from *Convolvulus Scammonia*, L

It is unfortunate that the name Jalapin should have been applied to the resin of *spurious* Jalap, which is identical with the *true* Resin of Scammony, and which is quite distinct from the official Resin of Jalap

During 1892 attention was again called to this misleading nomenclature (*P J* (3) xxii 888), and considerable correspondence ensued. It appears that it has been customary in this country to apply the term 'Jalapin' to the true Jalap Resin, but the article imported from Germany under that name is invariably the Ether soluble Resin from *spurious* Jalap or Scammony. Several suggestions were made, but none which seemed at all likely to be acceptable both in Britain and Germany. The most feasible proposal is that the term '*Scammonin*' should be used to designate the Ether soluble Resin (shown, *P J* (3) xxiii 86, to be identical from either of the previous named sources), and that the earliest opportunity should be taken to make official, under the name *Jalapin*, an Ether-wholly insoluble Resin from true Jalap

Not Official

JAMBUL

The Seeds of *Eugenia Jambolana*, Lam, which have been used in India and this country for reducing the amount of sugar excreted in diabetes, —*P J* (3) xviii 921, *B M J* '91, ii 1283, *B M J E* '92, i 894, *T G* '93, 611, *I J* li 138, *B M J* '01, ii 618

The dose should be large, 1 drm to 1 oz daily —*B M J* '91, ii 1284

Two cases in which 2 oz were given daily —*I J* li 139

It can also be given in the form of **fluid extract** (1 in 1) Dose, 10 to 60 minims = 0.6 to 3.6 cc

Not Official

JEQUIRITY

The Seeds of *Abrus precatorius*, L

Infusum Abri, 8 of the seeds to 100 of Water at 120° F, has been used in the treatment of granular lids, it sets up a purulent conjunctivitis, varying in

* Turpeth is the dried Root and Stem of *Ipomœa Turpethum* and is official in the *Ind* and *Col Add* for India and the Eastern and North American Colonies.

intensity with the strength and frequency of the A somewhat dangerous remedy. A very strong infusion, 1 to 4, was used by Dr Shoemaker in the treatment of affections of the skin—*Med. Bulletin*, Nov 1884, *L* '85, ii, 733, *L.M.R.* '86, 126, *T.G.* '87, 640, and *L.M.R.* '86, 541. Dr Martin's researches show that the determining causes of the inflammation and the toxic properties in general are due to a globulin and an albumose, the activity of which is rapidly destroyed by a moist heat of 85° C (185° F).

It has been shown that the continuous use of Abum produces tolerance to its toxicity—*B.M.J.* '97, ii 705

Foreign Pharmacopœias —Official in Spain

Jequiritol —A substance allied to Abum, supplied in sterile solution, containing 5 per cent. It possesses, when applied locally, a distinctly marked curative action on inflamed conjunctiva, when controlled by Jequiritol Serum, it is the best means for the removal of nebulae of the cornea—*L.* '01, i 1836

The root has been used in many hot countries for the same purpose as liquorice-root, hence it is called Indian Liquorice, but considering the known poisonous character of the seed the title is dangerously misleading.

The root and an extract prepared from it are official in the Pharmacopœia of India.

Not Official

JUGLANS

The Root-bark of *Juglans cinerea* Linn (Butternut), collected in autumn.

A mild cathartic, used in the form of **Extractum Juglandis**, prepared with Dilute Alcohol, dose, 5 to 10 grains = 0.32 to 0.65 gramme, and **Juglandin**, an eclectic remedy, used in doses of 5 to 10 grains.

Not now official in U.S.

FOLIA JUGLANDIS —The Leaves of *Juglans regia* L (Walnut) are Official in Austr., Belg., Ger., Mex. and Spain (Hoja de Nôgal), Belg. has also a fluid extract.

FLUIDEXTRACTUM JUGLANDIS —From the inner bark of the root. Made with Alcohol (49 p.c.), 1 cc. of fluid extract represents 1 gramme of drug—*U.S.N.F.* It is used as a cathartic. Average dose 1 fl. drm. = 3.6 cc.

SPIRITUS NUCIS JUGLANDIS —A distilled preparation from the Walnut (*Juglans Regia*).

Aromatic bitter, astringent.

Dose —1 to 4 fl. drm. = 2.6 to 14.2 cc.

JUNIPERI OLEUM.

OIL OF JUNIPER

FR., ESSENCE DE GENÉVRIER, GER., WACHOLDERÖL, ITAL.,
ESSENZA DI GINEPRO

A colourless, or pale yellow or yellowish-green oily liquid, having a characteristic odour, and balsamic, burning and somewhat bitter taste. It is the Volatile oil distilled from Fruit of *Juniperus communis* Linn. The Fruits should be full-grown and unripe.

It should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from air and light. The Oil has a tendency to resinify on keeping, and old Oil is more viscid, has an acrid reaction and has a somewhat rancid odour. The solubility is also affected, the Oil becoming less soluble in Alcohol (90 p.c.)

The Oil contains the terpene Pinene, boiling point 156°C (312°F), the sesquiterpene, Cadinene, boiling point 274°C (525°F) and Juniper Camphor, and an Ester, boiling point 180°C (356°F)

Empyreumatic Oil of Juniper is given under Cadmi Oilum, p 271

Solubility—1 in 20 of Alcohol (90 p c), but it does not become quite clear, it mixes with equal parts of Absolute Alcohol, but if more Alcohol be added it becomes milky

Medicinal Properties—Cumulative, antispasmodic, and a stimulating diuretic, the latter property constituting its chief medicinal value. Used in cardiac and hepatic diopysical cases, either alone or combined with other diuretics, should not be used in acute Bright's disease

Dose— $\frac{1}{2}$ to 3 minims = 0.03 to 0.15 c c

Official Preparation—Spiritus Juniperi, contained in Mistura Crocosoti

Not Official Spiritus Juniperi Compositus

Foreign Pharmacopœias—Official in Austri, Ger and Jap, sp gr 0.864 to 0.880, Norw, sp gr 0.860 to 0.870. Hung, sp gr 0.840 to 0.900, Fr (Genièvre) and Ital (Essenza di Ginepro), sp gr 0.865 to 0.895, Port (Essencia de Zimbro), sp gr 0.955 to 0.879, Swiss, sp gr 0.860 to 0.885, U S, sp gr 0.860 to 0.900 at 25°C (77°F)

Tests—Juniper Oil has a sp gr of 0.865 to 0.895, which figures are increased by exposure to air or by age. It is officially required to dissolve in 1 volume of a mixture consisting of equal parts of Absolute Alcohol and Alcohol (90 p c), but the solubility of the Oil also depends greatly upon the length of time it has been kept. The *BP* gives no figures for optical rotation. It is usually levogyrate, -3° to -12° in a tube of 100 mm. No indication is given in the *BP* as to the temperatures at which the various fractions of the Oil should distil. It is in the relative proportion of Pinene to Cadinene that English oils chiefly differ from foreign oils. An English oil examined by Bud (*C'D* '07, n 172) showed a 12 p c fraction distilling between 155° to 160°C (311° to 320°F), as against 37 p c fraction in foreign oil, a 35 p c fraction between 160° to 180°C (320° to 356°F) as against 34 p c fraction in the foreign oil, a 19 p c fraction between 180° to 255°C (356° to 491°F) as against a 10 p c fraction in the foreign oil, and a 22 p c fraction at 255° to 280°C (491° to 536°F), as against a 10 p c fraction in the foreign oil. It has been recommended (*C'D* '07, n 355) that the *BP* requirements with regard to fractionation should be, not less than 50 p c nor more than 60 p c, should distil below 165°C (329°F), that the refractive index of the unfractionated oil should not be less than 1.4750 (at 20°C (68°F)), and the residue after distilling 80 p c should have a refractive index at 20°C (68°F) of not less than 1.4900 nor more than 1.4950.

The more generally occurring sophistications are Turpentine Oil, Juniper Wood Oil and Alcohol. Of these, Alcohol is the only one readily detected. It is contained in the first fractions of the Oil, and is identified by the formation of Iodoform on warming this fraction

with Potassium Hydroxide Solution and sufficient Iodine Solution to ensure a slight excess Turpentine Oil is very difficult of detection except when present in large amount

Preparation

SPIRITUS JUNIPERI SPIRIT OF JUNIPER

Oil of Juniper, 1, Alcohol (90 p c), *q s* to yield 20 If not bright, filter through Tale (1 in 20)

Dose—20 to 60 minims = 1 2 to 3 6 c c

It is two and a half times stronger than *B P* '85

Foreign Pharmacopœias—Official in Jap, 1 in 50, by weight, U S, 1 in 20, Austr, Ger and Swiss, 1 fruit in 4, by distillation, Port and U S. have a compound spirit Not in the others

Not Official

SPIRITUS JUNIPERI COMPOSITUS—Oil of Juniper, 0 1, Oil of Caraway, 0 05, Oil of Fennel, 0 05, Alcohol (95 p c) 70, Water, *q s* to make 100—*U S P*

This has been incorporated in the *B P C* using 75 of Alcohol (90 p c)

Not Official

KALADANA

Syn—**PHARBITIS NIL**

The dried Seeds of *Ipomœa hederacea*, Jacq Cathartic, resembling Jalap in action

Official in the *Ind* and *Col Add* for India and the Eastern Colonies, as are also the **Compound powder**, Kaladana, 5, Acid Potassium Tartiate, 9, Ginger, 1, dose, 20 to 60 grains = 1 3 to 4 grammes, the **Tincture**, 1 of seeds in 5 of Alcohol (70 p c), dose, 30 to 60 minims = 1 8 to 3 6 c c, and the **Resin**, dose, 2 to 8 grains = 0 13 to 0 52 gramme

The Compound Powder, Tincture and Resin have been incorporated in the *B P C*

Not Official

KAMALA

Syn—**GLANDULÆ ROTTLERÆ**

A fine, granular mobile, brick-red powder, and hair, obtained from the surface of the Fruits of Arg

Solubility—Almost insoluble in Water, but about 60 p c of a sample (containing 6 p c of ash) was soluble in Absolute Alcohol, in Chloroform, and in Ether, and was for the most part soluble in Liquor Potassæ

Anthelmintic and purgative Successfully given in tænia, in doses in 30 to 120 grains = 2 to 8 grammes

Prescribing Notes—The powder is usually given suspended in Gruel, Mucilage, Juice or Syrup, or it may be prescribed as a dry mass In 1 lb of Male 1 lb of female seeds will make 1 lb of powder, it need be, for

Foreign Pharmacopœias—Official in Austr and Hung (10 p c of ash), Ger, Ital Jap, Swed and Swiss (6 p c of ash), Hung has also Kamala Depuratum, Port, and Russ (8 p c of ash), Mex Not in the others

TINCTURA KAMALÆ—Kamala, 1, Alcohol (60 p c), 5

Dose—1 to 2 fl drms = 3 6 to 7 1 c c.

KAOLINUM.

KAOLIN

NO Syn—CHINA CLAY, PORCELAIN CLAY

A native Aluminium Silicate, powdered, and from which the gritty particles have been removed by elutriation

A fine white clay, derived from the decomposition of the felspar of granitic rocks, extensive tracts of it occur in Cornwall. When finely ground and washed it is used as a form of Fuller's Earth

Has been used in Germany for many years as an excipient for pills of the easily reducible salts of metals, such as Gold Chloride, Silver Nitrate, and Potassium Permanganate, but a mixture of Paraffin answers better. See MASSA PARAFFINUM, p 863. It is also employed for clarifying Wine, Beer, and Syrups

Official Preparation—Contained in *Pilula Phosphori*

Not Official—*Cataplasma Kaolini*, *Unguentum Kaolini* and *Massa Kaolini*

Foreign Pharmacopœias—Official in Aust, Belg, Dutch, Ger, Hung, Jap and Swiss, (*Bolus Alba*), Dan, Norw, Swed and U.S. (*Kaolinum*) Not in the others

Tests—Kaolin, when fused with Potassium or Sodium Hydroxide or Carbonate, and the fused product, when cold treated with Water, yields a solution, which, neutralised with Hydrochloric Acid, affords a gelatinous precipitate of Silica, and if the liquid be evaporated to dryness, redissolved in Water, and filtered, the filtered liquid yields with Ammonia Solution a white gelatinous precipitate, insoluble in excess of the reagent, soluble in diluted Hydrochloric Acid. Another portion of the filtrate yields with either Potassium or Sodium Hydroxide Solution a white gelatinous precipitate, soluble in an excess of the reagent. The *USP* mixes the Kaolin with Water and Sulphuric Acid, evaporates the mixture to dryness, and heats the residue until fumes of Sulphuric Anhydride appear. When the residue is treated with boiling Water and filtered, a grey insoluble deposit of impure Silica remains on the filter. The addition of the Sulphuric Acid to the mixture of Kaolin and Water should cause no effervescence, indicating the absence of Carbonates. The *USP* includes a test for the absence of more than traces of Iron by mixing 2 grammes of the Kaolin with 10 cc of Water and one-quarter of its weight of Sodium Salicylate, and requires that not more than a slight reddish tint shall be produced. The non-volatile residue left on ignition at a red heat should amount to not less than 85 pc.

Not Official

CATAPLASMA KAOLINI—Kaolin, in very fine powder, 547, Boric Acid, 45, Thymol, $\frac{1}{2}$, Methyl Salicylate, 2, Oil of Peppermint, $\frac{1}{4}$, Glycerin, 375, all by weight, *USP*. Heat the Kaolin in a suitable vessel at 100° C (212° F) with occasional stirring for one hour, well mix with the Boric Acid, and then incorporate thoroughly with the Glycerin, finally add the Thymol which has been dissolved in the Methyl Salicylate and the Oil of Peppermint, and make a homogeneous mass which should be kept in an air tight container.

The *US Dispensatory* states that the quantity of Glycerin will be found insufficient for some kinds of kaolin, and the *BPC* has incorporated this form, making a slight change in the quantity of Kaolin and Glycerin as recommended.

Kaolin, 52 70, Boric Acid, 4 50, Thymol, 0 05, Methyl Salicylate, by weight, 0 20, Oil of Peppermint, by weight, 0 05, Glycerin, by weight, 42 50 same directions as above, *B P C*

UNGUENTUM KAOLINI—Soft Paraffin, 1, Hard Paraffin, 1, melt, and add Kaolin, 1, stir till cold

This has been proposed as a basis for pills containing Silver Nitrate or Potassium Permanganate—*P J* (3) xv 60

A very great improvement upon it is the following—

MASSA KAOLINI—Soft Paraffin, 2, Hard Paraffin (m p 120° F) 1, Kaolin, 1 This will make a good mass with three times its weight of Potassium Permanganate

A mixture of Hard Paraffin (m p 120° F), 1, with Soft Paraffin, 1½, answers even better, and will make a good mass with four times its weight of Permanganate See **MASSA PARAFFINUM**, p 863

Both of the above masses have been incorporated in the *B P C*

Not Official

KAVA-KAVA

The decorticated, dried and divided Root of *Piper Methysticum*, Forster. Spinal depressant, causing loss of muscular power, diuretic, used in chronic catarrhal conditions of the genito-urinary organs Used by the inhabitants of the Polynesian Isles in the preparation of an intoxicating liquor

Official in the Ind and Col Add for the Australian Colonies

An **Extract**, prepared with diluted Alcohol, used as a hypnotic, dose 1 to 5 grains = 0 06 to 0 32 gramme, also a **Fluid Extract** (1 in 1), dose 15 to 60 minims = 0 9 to 3 6 c.c.

An **Extractum Kavae Liquidum** (1 in 1), dose 30 to 60 minims = 1 8 to 3 6 c.c., is official in the *Ind and Col Add* for the Australian Colonies

Not Official

KERATIN

A substance introduced by Dr Unna for coating pills which are intended to pass the stomach and act in the small intestine It is made by digesting horn shavings, first in artificial gastric juice (acidified Pepsin solution) until all the albuminous substances have been dissolved, and treating the residue with Ammoniacal Solution The ammoniacal solution, when evaporated, yields a gum-like liquid, which can be used for coating pills The coating, although unaffected by Hydrochloric Acid, is soluble to some extent in Acetic and Citric Acids, which should therefore not be given at the same time

LIQUOR KERATINI—Keratin, 1, Alcohol (90 p.c.), 5, Strong Solution of Ammonia, 5, Alcohol and Ammonia and dissolve the Keratin

This has been incorporated in the *B P C*

This makes a thin coating, and dries quickly It is better to give the pills a thin coating of Oil of Peppermint, two coatings of Keratin, and then a coating of Oil of Peppermint

KINO.

KINO

The juice obtained from incisions in the trunk of *Picrostaphis Marsupium*, Roxb, evaporated to dryness See Descriptive Notes below

Medicinal Properties—A powerful astringent Employed in obstinate diarrhoea and dysentery in the form of compound powder or with chalk, also in passive hæmorrhage Externally as a styptic

Dose, in powder, 5 to 20 grains = 0.32 to 1.3 grammes

Prescribing Notes—Generally given in the form of the compound powder, it may be administered in the form of cachets The Tincture is useful in gargles and tooth washes, the Lozenges for throat affections

Incompatibles—Mineral Acids, Alkalis and Carbonates, Metallic salts, and Glycerin

Official Preparations—Pulvis Kino Compositus and Tinctura Kino Contained in Pulvis Catechu Compositus

Not Official—Trochisci Kino

Foreign Pharmacopœias Official in Jap, Port, Swiss and U S Not in the others

Descriptive Notes The official Kino is distinguished in commerce as East Indian, Malabar, or Cochin Kino, it being often shipped from that port It consists of the juice of the tree dried without artificial heat, but the official article is stated to be the juice obtained from incisions in the trunk of *Pterocarpus Marsupium* evaporated to dryness But if this implies artificial evaporation, such a drug is not a commercial article The official drug is in small angular, glistening, reddish black, brittle, opaque fragments, which in thin splinters have transparent and ruby red edges It has no odour, is very astringent, and when chewed tinges the saliva red

Tests—Kino is officially stated to be partially soluble in cold Water, and that not less than 80 p.c. should be soluble in boiling Water Only 88 grains out of 100 grains of Tellicherry Kino are dissolved by cold Water, and 35 grains of Isinglass will precipitate the whole of the astringent matter from the solution Compared with Pale Catechu it is more soluble in Water, and the solution is more astringent The *USP* says slowly soluble in cold Water The *BP* states that it is almost entirely soluble in Alcohol (90 p.c.), and yields little or nothing to Ether, the *USP* that it is soluble in Alcohol (94.9 p.c.), and nearly insoluble in Ether The ash varies from 1 to 2 p.c.

Kino Eucalypti (dose, 5 to 20 grains = 0.32 to 1.3 grammes) is official in the *Ind* and *Col Add* for the Australian Colonies

Butea Gum (Bengal or Madras Kino), the assayed juice from the stem of *Butea frondosa*, Roxb., is made official in the *Ind* and *Col Add* for use in place of Kino in India and the Eastern Colonies Kino Eucalypti and Butea Gum have the same uses and doses as East Indian Kino It becomes insoluble if long kept

Butea Seeds and Powder of the same are also included for the same countries as an anthelmintic for ascariæ, 10 to 20 grains of the Powder

Warning (*Ph Ind*) gives the dose as 20 grains three times a day for three days, and a dose of Castor Oil on the fourth day, but its use requires care

Preparations

PULVIS KINO COMPOSITUS COMPOUND POWDER OF KINO
Kino, 15, Opium, 1, Cinnamon Bark, 4 (1 Opium in 20)
Keep it in a well closed vessel

Dose $\frac{1}{2}$ to 20 grains = 0.32 to 1.3 gramme

TINCTURA KINO. TINCTURE OF KINO

Kino, in powder, 2, Glycerin, 3, Distilled Water, 5, Alcohol (90 p c), q s to yield 20

Dose $\frac{1}{2}$ to 1 fl dm = 1.8 to 3.6 c c

Foreign Pharmacopœias—Official in Swiss, 1 in 5, by weight, U S, 1 in 20, see below Not in the others

Tests—Tincture of Kino has a sp gr of 0.988 to 0.990, contains from 22.0 to 25.0 p c w/v of total solids, and about 45 p c w/v of Absolute Alcohol. A standard of 5 p c w/v of Kinotannic Acid has been proposed (C D '98, n 971) for the Tincture

As gelatinisation is probably due to an enzyme, the following formula has been proposed: Kino, 2 oz., Boiling Water, 10 fl oz. Add the Kino to the Water in a suitable vessel, and maintain the whole at or near the temperature of 100° C for fifteen minutes, agitating frequently. Allow to cool, replace the Water lost by evaporation, add Alcohol (90 p c) 10 fl oz, and set aside for twelve hours, then strain.—P J '08, n 702

Rub 5 of Kino and 1 of Purified Talc with 15 of Glycerin and 20 of Distilled Water, transfer to a flask and weigh, heat it on a water-bath for one hour and, after cooling, add Water to make up any loss, then add 65 of Alcohol (95 p c), mix and filter through Purified Cotton, and pass through the cotton Alcohol (95 p c) q s to make 100.—U S P

Not Official

TROCHISCI KINO—Containing 2 grains in each lozenge, with Black Currant paste

This has been incorporated in the B P C.

Not Official

KOLA

The Seeds of *Cola acuminata*, Schott and Endl, a tree whose habitat is the Western Coast of Africa, between Sierra Leone and the Congo. The Seeds contain 2 to 2.5 p c of Caffeine, to which it owes its virtues, also a glucoside Kolanin. Kola is official in the *Fr Codex* (1908) and is required to contain at least 1.25 p c of Caffeine, an Extract which is required to contain at least 10 p c of Caffeine and 1 Fluid Extract, which is required to contain at least 1.25 p c of Caffeine. Caffeine is also official. A proprietary article is also sold under this name. Various preparations have been made, *viz*, Kola-chocolate, Kola elixir, Kola wafers, Kola wine, also Fluid Extract

Exerts an action in cases of fatigue on the natives of those parts of Africa where it grows, whereas preparations of Kola made in Europe from the dried nuts are much less active. The fresh nuts and the extract obtained therefrom contain a phenolic compound, Kolatone, which exerts an action entirely different from that of Caffeine, to which the medicinal properties of Kola in fatigue have hitherto been ascribed.—L '06, n 177

Teinture de Kola (1 to 5 of Alcohol 60 p c) is official in Fr, Swiss (Extractum Colæ Fluidum) contains 1.5 p c of Caffeine and Theobromine, also (Vino de Kola) 1 of Fluid Extract and 95 of Vinum Meridianum, Austerum, Spain, Alcohol 70 p c (Tintura Alcoholica de Kola), 1 Kola in 10 with Alcohol 70 p c, and (Vino de Kola) 1 of Kola in 10 of Sherry. All by weight

B P C has an Extractum Kolæ Liquidum, 1 in 1 using Alcohol (60 p c), Elixir Kolæ, 1 of Liquid Extract, $\frac{1}{2}$ of Vanilla and Syrup to produce 100, also Vinum Kolæ, 1 of Elixir of Kola and Detannated Sherry to produce 8.

Descriptive Notes—Kola nuts are imported from West Africa, and to some extent from the West Indies. The so called nuts consist of seeds freed from the seed coats, and are sometimes broken up into two cotyledons or into four. These are derived, according to Schumann, from two different species, those having two cotyledons, carmine red when fresh, from *Cola vera*, Schum., and those with four cotyledons, sometimes red and sometimes white when fresh, from *C. acuminata*. The seeds of *C. lupulata*, Schum., are also used by the natives but there is no evidence that they are exported. The seeds as imported are about $1\frac{1}{2}$ inch (37 mm) long and $1\frac{1}{4}$ inch (31 mm) in diameter, irregularly ovate oblong, with a more or less oblique line where the two cotyledons meet, but those imported from Trinidad and the West Indies are often not much more than half this size. Kola seeds are hard, solid, tough, and of a reddish brown colour. The taste is earthy and somewhat astringent and slightly bitter.

KOUSSO. *See* CUSSO.

KRAMERIÆ RADIX.

KRAMERIA ROOT

B P *Syn*—RHATANIA ROOT

FR, RATANIA DU PIROU, GER, RATANIARWURZEL, ITAL, RATANIA,
SIAN, RATANIA

The dried Root of Para Rhatany, a species of *Krameria*, attributed to *Krameria argentea*, Mart. or of (2) Peruvian Rhatany, *Krameria triandra*, R. and P.

Medicinal Properties—A powerful astringent, tonic. Used in chronic diarrhœa, in passive hæmorrhages and mucous discharges, as menorrhagia and leucorrhœa, and generally where Tannin and Catechu are beneficial. The infusion is used as a gargle in relaxed sore throat, one teaspoonful of the tincture in a wineglassful of water is an excellent wash for spongy and inflamed gums, or stomatitis due to Mercury. Locally, in form of suppository with Opium or Morphine, it is used in prolapsus ani, anal fissure, and bleeding piles.

Dose—20 to 60 grains = 1.3 to 4 grammes, in powder.

Incompatibles—Alkalis, Lime Water, Iron and Lead salts, Gelatin.

Official Preparations—Extractum *Krameriæ*, Infusum *Krameriæ*, Liquor *Krameriæ* Concentratus, Tinctura *Krameriæ*, Trochiscus *Krameriæ* and Trochiscus *Krameriæ* et Cocaine. Contained in Pulvis Catechu Compositus.

Not Official—Extractum *Krameriæ* Fluidum, Gossypium *Krameriæ*, Infusum *Krameriæ* Concentratum, Suppositorium *Krameriæ*, Sympus *Krameriæ* and Trochiscus *Krameriæ* et Boracis.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Jap. (Rhatany Root), Norw., Russ., Swed. and Swiss (Ratanhia), Mex. (*Krameria*), Hung. (Ratanhia), Ital., Port. and Span. (Ratania); U.S. (*Krameria*).

Descriptive Notes—The roots of two species are official in the B.P. Peruvian Rhatany, the root of *Krameria triandra*, Ruiz

and Pavon, is 1 - - - 1 by having a large woody crown, often 2 or more in in diameter, giving off several tapering cylindrical roots. These are tough and not easily broken, of a dark brownish-red colour, the rough bark having a splintery fracture, and readily separating from the woody centre, which is of a yellowish colour. Para Rhatany is derived from *Krameria argentea*, Martius, it consists of cylindrical roots, 12 to 18 in long and $\frac{1}{4}$ to $\frac{1}{2}$ in in diameter, has a purplish-brown colour, and smooth thick bark, cracked transversely at intervals, with a short fracture, and adhering closely to the reddish-brown wood. The bark of both kinds is very astringent, and when chewed the root tinges the saliva red, but the woody portion is almost tasteless. The powder of Peruvian Rhatany is lighter in colour than that of the Para drug, and is characterised by conical or pear-shaped starch grains, often arranged in a stellate form or in groups of three or four, by the flattened bast fibres, by the prismatic crystals of Calcium Oxalate in the bast parenchyma, and by the arrangement of bast fibres which, instead of forming extended groups, are dispersed irregularly. Formerly a variety known as Savanilla Rhatany was imported, resembling the Para in appearance, but of a paler purplish tint, and thicker bark, being one-third to a quarter of the thickness of the wood. It can be distinguished by thin sections of the root giving a violet colour when moistened with a ferrous salt: those of the Peruvian sort are more greyish hue, and those of Para bluish-black. Savanilla Rhatany is derived from *Krameria tomentosa*, St Hil. A so-called Rhatany Root from Guayaquil was offered in commerce a few years ago. It contains cluster crystals of Calcium Oxalate, which do not occur in the other kinds of Rhatany mentioned above.

Tests - Krameria Root contains from 1 to 2 p.c. of ash

Preparations

EXTRACTUM KRAMERIÆ EXTRACT OF KRAMERIA *B.P. Syn.*
—EXTRACT OF RHATANY

Prepared from Krameria Root, by exhaustion with Distilled Water and evaporation to dryness.

Dose — 5 to 15 grains = 0.32 to 1 gramme

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Dutch, Fr., Jap., Mex., Port., Russ., Spain and U.S., Hung., crude Extract purified with warm Water, Ital., U.S., made with boiling Water, Belg., Mex., and U.S. have also a Fluid. Extract. Not in Ger. or Norw.

INFUSUM KRAMERIÆ INFUSION OF KRAMERIA. *B.P. Syn.*—
INFUSION OF RHATANY

Krameria Root bruised, 1, boiling Distilled Water, 20 Infuse 15 minutes. (1 in 20)

Dose — $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 c.c.

This Infusion should be freshly prepared, as it deposits when kept.

Foreign Pharmacopœias.—Official in Fr. and Mex., Tisane, 1 in 50. Not in the others.

LIQUOR KRAMERIAE CONCENTRATUS CONCENTRATED SOLUTION OF KRAMERIA

10 of Krameria Root, in No 40 powder, percolated with Alcohol (20 p c), to yield 20 (1 in 2)

Dose — $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

Tests — Concentrated Solution of Krameria has a sp gr from 1 010 to 1 020, contains about 10 p c w/v of total solids, and about 18 p c w/v of Absolute Alcohol

TINCTURA KRAMERIAE TINCTURE OF KRAMERIA *B P Syn*
TINCTURE OF RHATANIA

4 Krameria Root, in No 40 powder, percolated with Alcohol (60 p c), to yield 20 (1 in 5)

In *B P* 1885 it was 1 in 4

Dose — $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

Foreign Pharmacopœias — Official in Aust, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Swed, Swiss and U S, 1 in 5 All by weight except U S — Not in Span

Tests Krameria Tincture has a sp gr of 0 930 to 0 940, contains about 5 p c w/v of total solids, and about 56 p c w/v of Absolute Alcohol

TROCHISCUS KRAMERIAE KRAMERIA LOZENGE *B P Syn* — RHATANIA LOZENGE

1 gram of Extract of Krameria in each, with Fruit Basis

Foreign Pharmacopœias — Official in U S — Not in the others

TROCHISCUS KRAMERIAE ET COCAINAE KRAMERIA AND COCAINE LOZENGE *B P Syn* — RHATANIA AND COCAINE LOZENGE

1 grain of Extract of Krameria and $\frac{1}{10}$ grain of Cocaine Hydrochloride in each, with Fruit Basis

Not Official

EXTRACTUM KRAMERIAE FLUIDUM — Krameria, in No 30 powder, 100, Glycerin, 10, Diluted Alcohol (49 p c), *q s* to produce 100 — *U S P* 1890. This has been incorporated in the *B P C* using Alcohol (15 p c)

GOSSYPIUM KRAMERIAE — Tincture of Rhatany, $\frac{1}{2}$ fl oz, Glycerin, 10 minims, mix and with it saturate evenly Cotton Wool, 60 grains, and dry.

INFUSUM KRAMERIAE CONCENTRATUM — Krameria, in No 40 powder, 40, Alcohol (90 p c), 25, Dilute Chloroform Water (1 to 1000), *q s* to make 100. Prepare by repercolation before the addition of the Alcohol to the reserved portion, this should be heated to a temperature of not less than 55° C and maintained therefor 5 minutes. Dose — 30 to 60 minims = 1 8 to 3 6 c c *Farr and Wright, P J '06, 1 165, and '07, 1 621, C D '06, 1 252, 1 B P 1907, 250*

This appears in the *B P C*

SUPPOSITORIUM KRAMERIAE — Extract of Rhatany, 8 grains, Morphine Hydrochloride, $\frac{1}{10}$ grain, Stearin, 10 grains

Foreign Pharmacopœias — Official in Fr and Span, 1 gramme of Extract = 15 5 grains in each

SYRUPUS KRAMERIAE Fluid Extract of Krameria, 45, Syrup, 55 — *U S P*

This has been incorporated in the *B P C*.

Foreign Pharmacopœias—Official in Belg, Fluid Extract 1, Syrup 9; Fr, Extract 25, Syrup 975, concentrate to 1000 by weight, Ital, Extract 2, Water 5, Syrup 98, concentrate to 100 by weight, Mex, Extract 25, Syrup 975, Swiss, 1 of Extract in 100

Not Official

LACHNANTHES TINCTORIA

A North American plant, known colloquially as Red Root or Spirit Weed, a native of the United States. A homœopathic remedy for checking the cough of phthisis, and for treating pneumonia and typhus. Did not exert any inhibitory action on the growth of tuberculosis, but rather seemed to hasten it.—*B M* '01, 11 747, 912, 1124, 1606, 1874, '02, 1 59, 101, 113, '02, 11 146, *L* '01, 11. 1605, '02, 11 72, *Pr* lxvii 493, *P J* '02, 1 103

Dose—2 to 10 minims of a 1 to 10 Tincture made with Alcohol (45 p c)
A solid and a fluid extract are also known commercially

Not Official

LACTUCA

Lettuce is the Flowering Herb of the wild indigenous plant, *Lactuca virosa*, L.

Medicinal Properties—A sedative in irritable cough, either in the form of Extractum Lactucæ or as Lactucarium

Dose—5 to 15 grains = 0.32 to 1 gramme

LACTUCARIUM—The juice from the incised flower-stalk of *Lactuca virosa* and other species, collected and dried

Dose—2 to 6 grains = 0.13 to 0.40 gramme

Foreign Pharmacopœias—Official in Dutch, Hung, Mex, Port, Spain and U. S. Not in the others

SYRUPUS LACTUCARIUM—Tincture of Lactucarium, 10, Glycerin, 20, Citric Acid 0.1, Orange-Flower Water (*U S P*), 5, Syrup, *qs* to produce 100. Mix the Tincture with the Citric Acid, add the Orange-Flower Water in which the Citric Acid has been previously dissolved, filter if necessary, add the Syrup and make up to 100.—*U S P*

Average Dose—2 fl drms = 7.1 c c

This has been incorporated in the *B P C*

TINCTURA LACTUCARII—Lactucarium, 50, Glycerin, 25, Alcohol (95 p c), Purified Petroleum Benzin, Diluted Alcohol (49 p c), Water, of each a sufficient quantity to make 100. Powder the Lactucarium with coarse sand and add 200 of Purified Petroleum Benzin, after macerating for 48 hours with frequent agitation pour the mixture on a double filter and allow to drain, wash the residue by gradually adding 150 of Purified Petroleum Benzin, and let the Lactucarium dry by exposure to air. Powder the dried Lactucarium using more sand if necessary, and pack it moderately in a percolator, mix the Glycerin with 20 of Water and 50 of Alcohol (95 p c), and moisten the powder with 50 of the mixture and macerate for 24 hours, then let the percolation proceed slowly, gradually adding first the remainder of the menstruum, and then Diluted Alcohol (49 p c) until the Lactucarium is exhausted, reserve the first 75 of percolate, evaporate the remainder, mix with the reserved portion, filter, and make up to 100 with Diluted Alcohol (49 p c).—*U S P*

Dose— $\frac{1}{2}$ to 1 fl drms = 1.8 to 3.6 c c

This has been incorporated in the *B P C*.

TINCTURA LACTUCARII ET OPII—Official in Mex. Alcoholic Extract of Lactucarium 2, Extract of Opium 1, Alcohol (50 p c), $q\ s$ to 80

SYRUPUS LACTUARII ET OPII—Official in Mex. 1 of the above Tincture in 50

TROCHISCI LACTUÆ—Lozenges containing 1 grain Extract Lactuce in each

LANOLIN. *See* ADEPS LANE

Not Official

LARICIS CORTEX

LARCH BARK

The Bark of *Larix Laricina*, DC., collected in the spring, deprived of its outer portion and dried. It contains a volatile crystallisable acid, **Larixinic Acid**, which sublimes in vapour of water.

Medicinal Properties—Similar to those of Oil of Turpentine. Useful in chronic bronchitis to diminish excessive secretion, the Tincture well diluted forms an astringent injection.

TINCTURA LARICIS—Tinch Bark, 1, Alcohol (90 p c), $q\ s$ to make 8 (1 in 8)

Dose—20 to 30 minims = 1 2 to 1 5 c c

This has been incorporated in the *L. P. C.*

TEREBINTHINA VENETA or **T. LARICIS** (Venice Turpentine)—A viscid liquid of a yellowish or greenish yellow colour, obtained from *Larix Europæa*, DC. It does not readily harden on exposure to air, or when mixed with $\frac{1}{8}$ of Magnesia. Soluble in absolute Alcohol. It is much used on the Continent, and in veterinary practice in this country.

Foreign Pharmacopœias—Official in Belg., Dutch, Fr. (Térébenthine du Molène), Hung., Ital. (Trementina di Venezia), Norw., Port., Russ., Span. (Trementina de Aleice), Swed. and Swiss. Not in the others.

VASOLIMENTUM TEREBINTHINÆ—Venice Turpentine, 20, Liquid Vasoliment, 80—*Hager*

PAROGENUM TEREBINTHINÆ—Venice Turpentine, factitious, 20, Parogen, $q\ s$ to produce 100—*L. P. C.*

LAUROCERASI FOLIA.

(CHERRY LAUREL LEAVES)

FR., LAURIER CERISER, GER., KIRSCHLOHRN FEH., ITAL., LAUROCERASO,
SPAN., LAUREL CERIZOS

Descriptive Notes—The fresh leaves of *Prunus Laurocerasus*, L., are official. There are about six varieties of the plant in cultivation in this country. The strongest and hardest is the variety *Caucasica*, which has darker green, thicker and less rounded leaves than the variety *Colehica*, in which they are more oblongate, more delicate and paler; the variety *Schiphaensis* has smaller leaves, about the size of those of the bay tree, and forms a small shrub only 3 to 5 feet high. M. Perrinelle is of opinion that the variety *Caucasica* should be official,

P T (3) *VIM* 170 The Leaves in all the varieties are coriaceous, shining above, but paler beneath where the midrib is prominent, and at the base on each side of the midrib there are one or two glandular depressions. The Leaves are lanceolate oblong, or more or less obovate oblong, 5 to 7 in long ($12\frac{1}{2}$ to 17 cm., *P B*) and $1\frac{1}{2}$ to 2 in wide, attenuated towards either end, or rounded in some varieties, with a slightly revolute margin, with short sharp serratures, and glandular teeth, which are more distant towards the base. The taste is astringent and bitterish. The Leaves are odorless until bruised, but then immediately emit an odour of bitter almonds and Hydrocyanic Acid, the Leaves can be dried whole and powdered, and still yield Hydrocyanic Acid when the powder is moistened. The glucoside yielding Laurocerasin occurs in the parenchyma of the leaf, and the emulsion in the endodermis of the veins. The Leaves contain the largest amount of Hydrocyanic Acid in July, and the least in February, as much as 2½ p c having been found in June in young leaves. Cherry-Laurel Water is liable to vary in percentage of Hydrocyanic Acid, according to the time of year that the Leaves are collected, whether the Leaves are young or fully matured, and also according to the variety employed, the finely-chopped Leaves also yield more Hydrocyanic Acid than if merely bruised.

Official Preparation—Aqua Laurocerasi

Foreign Pharmacopœias Official in Belg., Dutch, Fr. (Laurier Cerise), Ital. (Lauroceraso), Port. (Loureiro-Gelejeira), and Span. Not in the others.

Dutch has an Oleum Laurocerasi

Preparation

AQUA LAUROCERASI CHERRY-LAUREL WATER

Fresh Cherry-Laurel Leaves, 16, Water, 50, distil 20, and standardise the distillate to contain 1½ p c of Hydrocyanic Acid, HCN.

Note—To ascertain if it lost much of its strength by keeping, a sample was taken which contained 0.104 p c, and placed in a pint bottle about three-quarters full for a month, it then gave 0.094 p c, the bottle was then kept for a week with only 3 oz in it, and then gave 0.093 p c, the same was then kept three days with the cork out, and then gave 0.038 p c.

It would appear therefore that when kept in a closed vessel, the sample is stable, but notwithstanding the adoption of an official standard, other samples will be found sometimes as low as half the official strength.

Medicinal Properties—Nervine sedative. Similar to Hydrocyanic Acid, but without the nauseous odour of the Acid. Used as a lotion to allay itching in cutaneous diseases, also as an adjunct to eye lotions (1 or 2 in 16).

Dose.—½ to 2 fl drms = 1.8 to 7.1 cc

20 minims = 1 minim Diluted Hydrocyanic Acid

Incompatibles—Same as Hydrocyanic Acid

Antidotes—In case of overdose, the antidotes should be as directed under Acidum Hydrocyanicum Dilutum, p. 54.

Foreign Pharmacopœias—Official in Aust., Belg., Dutch, Ital., Span. and Swiss 1.0 HCN per 1000, Fr., 55 to 7 per 1000, Port., Leaves 1 in 2, not standardised. Not in the others.

The Brussels Commission of 1856 fixed 1 per 1000.

Tests—Cherry Laurel Water is officially required to contain 0.1 p.c. of absolute Hydrogen Cyanide. The method adopted by the *B.P.* for the determination is a volumetric one, and is described in the large type under 'Acidum Hydrocyanicum Dilutum'. It is to be assumed that a quantity proportionate to the difference in strengths between the two preparations is to be employed, but no statement to this effect appears. The comments made on the process appearing in the large type apply also here. The alternative process there suggested may be employed for this assay, using 50 c.c. of the Water.

LAVANDULÆ OLEUM.

OIL OF LAVENDER

FR., ESSENCE DE LAVANDE, CHÈRE, LAVENDULIER, TIAR, L'ESSENZA DI LAVANDA, SIAN, ESSENCIA DE ESSENGO

A pale yellow, or yellowish green, oily liquid, having a pleasant characteristic odour, and an aromatic and somewhat bitter taste. It is the volatile Oil distilled from the Flowers of *Lavandula vera*, DC.

It should be kept in well closed glass bottles of dark amber tint in a cool atmosphere, and it should be protected as far as possible from the light.

The principal constituents are an Alcohol Linalool $C_{10}H_{18}O$, eq. 152.98, identical with that obtained from Lignum Aloes, and its Acetic Ester (Linalyl Acetate), which also forms the principal constituent of Oil of Bergamot.

It contains also the terpenes, Pinene and Limonene, a second Alcohol, Geraniol, and a sesquiterpene. English Oil of Lavender contains from 7 to 10 p.c. of Esters calculated as Linalyl Acetate ($C_{12}H_{20}O$, $C_{10}H_{18}O$, eq. 194.68), whilst the French Oils contain from 25 to 50 p.c. Cineol is also present to a greater extent in the English than in the foreign oils. Gildemeister and Hoffmann state that the value of Lavender Oil depends on its content of Linalyl Acetate, but Parry, in common with most others, is of opinion that no comparison between the oils can be made on the basis of their Ester content, Linalyl Acetate not being the sole odouriferous constituent of Lavender Oil. Coumarin has been detected in French Lavender Oil.

It is sometimes adulterated with the foreign Oil of *L. vera*, DC., and the foreign Oil is frequently adulterated with Oil of Spike from *L. spica*, DC. The flavour is stated to be improved by keeping for a year after distillation, and then mixing with an equal volume of Absolute Alcohol.

Solubility—In all proportions of Alcohol (90 p.c.) and Absolute Alcohol, sparingly soluble in Alcohol (60 p.c.).

Medicinal Properties—An aromatic, gastric stimulant and carminative. Useful in flatulence and colic.

Dose.— $\frac{1}{2}$ to 3 minims = 0.03 to 0.12 c.c.

Prescribing Notes—The oil is rarely given alone, it is used as an adjuvant to other medicines. Small doses of the spirit are given on Sugar. The Compound Tincture is a favourite colouring for mixtures.

Official Preparations—Of the Oil, Spiritus Lavandulae, and Tinctura Lavandulae Composita Contained in Linimentum Cumphora Ammoniatum The Compound Tincture is contained in Liquor Aiscenicis

Foreign Pharmacopœias.—Official in Austr., Belg., Dan., Ger., Norw. (Ætheroleum Lavandulae), Ital. (Essenza di Lavanda), Russ., and Swed., sp gr 0 885 to 0 895, Dutch, sp gr 0 880 to 0 890, Fr., sp gr 0 882 to 0 895, Hung. and Jap., sp gr 0 885 to 0 900, Poit. (Essencia de Alfazema), sp gr 0 875 to 0 940, Spun. (Essencia de Espilego), sp gr 0 87 to 0 94, Swiss, sp gr 0 882 to 0 895, U.S., sp gr 0 880 to 0 892 at 25° C (77° F)

Tests.—Lavender Oil has a sp gr of 0 885 to 0 895, the official requirement is not below 0 885, the *USP* 0 875 to 0 910 at 25° C (77° F), the *PG* 0 885 to 0 895 The optical rotation in a tube of 100 mm is from -3° to -10° It is officially required to dissolve in 3 times its volume of Alcohol (70 p c), which corresponds with the requirements of the *USP* The *PG* specifies Alcohol (68 to 69 p c) and the parts refer to parts by weight All specimens of Lavender Oil will not yield a clear solution with 3 to 3½ volumes of the weaker strength Alcohol of the *PG* Neither the *BP* nor the *USP* includes a process for the determination of the Ester content The *PG* volumetric determination expressed in terms of Linalyl Acetate indicates at least 29 5 p c A standard of not more than 11 p c of Linalyl Esters has been suggested (*YPB* '03, 248) and not less than 36 p c for the French Oils

The more generally occurring sophistications are Turpentine Oil, Spike Oil, Spanish Lavender Oil, Rosemary Oil, and Alcohol Turpentine Oil lowers the sp gr and effects the rotation according to whether the Turpentine Oil used as the adulterant is dextrorotatory, *e.g.*, American Turpentine Oil, or levorotatory, *e.g.*, French Turpentine Oil Spike Oil increases the sp gr and lowers the rotation Spanish Lavender Oil behaves in a similar manner Rosemary Oil increases the sp gr and lowers the rotation, but renders the Oil less rotatory

Various substances, *e.g.* Ethyl Succinate, Benzoic Acid, and Salicylic Acid, have from time to time been added to the oil with a view to masking adulteration by artificially raising the Ester content The presence of Ethyl Succinate and Benzoic and Oxalic Acids may be determined by saponifying a weighed quantity of 2 grammes of the Oil with Potassium Hydroxide Solution, or with Acetic Acid, diluting to 50 c c and adding 10 c c of saturated Barium Chloride Solution The mixture is warmed for 2 hours on the water-bath and allowed to cool The formation of a crystalline precipitate indicates adulteration Salicylic Acid may be detected by the purple-violet coloration produced on the addition of Ferric Chloride TS after saponification and by the m p of the isolated acid The presence of Benzoic Acid may be confirmed by Ferric Chloride TS and the m p of the isolated acid The presence of Alcohol may be determined by a diminution in volume when the Oil is shaken with Water

Adulteration with Glycerin Monacetate has also been noticed Such adulteration may be detected by shaking the suspected oil with 4 to 5 volumes of Petroleum Ether, in which the Glycerin ester is insoluble, and Glycerin may be further identified by the Acrolein reaction.

Water—When the Oil is shaken with Water in a narrow graduated cylinder its volume should not be diminished (absence of Alcohol), *U S P*

Volumetric Determination—If 1 gramme of the Oil be heated in a reflux condenser with 10 cc of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution for half an hour on a water bath, and after cooling and the addition of a few drops of Phenolphthalein T S, the mixture be titrated with Semi-normal Volumetric Hydrochloric Acid Solution, at most 7 cc of the Acid should be necessary to discharge the colour, *P G*

Preparations

SPIRITUS LAVANDULÆ. SPIRIT OF LAVENDER

Oil of Lavender, 1 Alcohol (90 p c), *q s* to yield 10 (1 in 10)

Dose—5 to 20 minims = 0.3 to 1.2 cc

It is 5 times the strength of *B P* '85

Foreign Pharmacopœias—Official in Russ., 1 in 100, Dan and Swed., 2 in 100, Jap., 3 in 100, U S., 5 in 100 all with the Oil, and all by weight except U S. Austri., Dutch, Ger., Ital. and Swiss, all 1 in 1, and Port 1 in 2, from the flowers. Not in the others.

Tests Spirit of Lavender has a sp gr of 0.835 to 0.838, and contains about 68 p c w/v of Absolute Alcohol

TINCTURA LAVANDULÆ COMPOSITA. COMPOUND TINCTURE OF LAVENDER

Oil of Lavender, 45 minims, Oil of Rosemary, 5 minims, Cinnamon Bark, bruised, 75 grains, Nutmeg, bruised, 75 grains, Red Sanders Wood, 150 grains, Alcohol (90 p c), 20 fl oz. By maceration, adding the Oils at the finish.

Dose— $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 cc

Tests.—Compound Tincture of Lavender has a sp gr of 0.835 to 0.840, contains about 0.5 p c w/v of total solids and about 88.0 p c w/v of Absolute Alcohol

Foreign Pharmacopœias—Official in Swed., Tinctura Lavandulæ aromatica, Oil of Lavender 2, Oil of Rosemary 1, Saigon Cinnamon 8, Red Sanders Wood 8, Myristica 8, Alcohol (64 p c) 300 U S., Tinctura Lavandulæ Compositum, Oil of Lavender Flowers 8, Oil of Rosemary 2, Saigon Cinnamon 20, Cloves 5, Myristica 10, Red Sanders 10, Alcohol and Water, each, a sufficient quantity to make 1000 Jap., Tinctura Lavandulæ Compositum similar to U S

Not Official

LEPTANDRIN.

CULVERS ROOT

The Rhizome and Rootlets of *Trionia ligularis*, L.

A cathartic, and stimulates the flow of bile

An Alcoholic **Extract**, average dose, 0.25 gramme (1 grain), and **Fluid Extract** (1 in 1), average dose, 3 cc (15 minims), each both official in U S

Leptandrin An eclectic remedy, used as an alternative, $\frac{1}{2}$ to $\frac{3}{4}$ grain = 0.016 to 0.032 gramme, as a purgative 2 to 4 grains = 0.13 to 0.26 gramme

LIMONIS CORTEX.

LEMON PEEL

FR., ECORCE DE CITRON, GER., CITRONENSCHALE, ITAL., SCORZA DI LIMONE, SPAN., LIMON

The fresh outer part of the pericarp of the Fruit of *Citrus Medica*, L., var *Limonum*, Hook &

Commercially the peel is cut in December, and is more readily obtainable at that time

Ind and Col Add authorises the use of Dried Lemon Peel in India and the Colonies, when Fresh Lemon Peel is not obtainable

Medicinal Properties—Bitter stomachic and tonic Added to stomachic medicines Chiefly used, however, to impart flavour to other medicines

Official Preparations Of the peel, Oleum Limonis, Symplicium and Tinctura Limonis Used in the preparation of Infusum positum and Infusum Gentiane Compositum The oil is contained in Laminentum Polasii Iodidi cum Saponi, Spiritus Ammoniac Aromaticus, Tinctura Guaiaci Ammoniacata and Tinctura Valerianae Ammoniacata

Not Official—Terpeness Oil of Lemon, Citral

Foreign Pharmacopœias Official in Fr (Citron), Ger, Hung, Ital (Cedro and Limon), Jap, Port (Limão), Russ, Swiss and US Not in the others

Descriptive Notes—Only the fresh peel is official On its inner surface there should be only a small amount of the white spongy portion of the rind In the P (the rind, separated in strips and dried, is official The larger rough, thick-skinned peel of *Citrus Medica*, L., the citron or cedrat of the French, are sometimes sold as lemons, but are rather less fragrant and contain less juice. These are usually imported in July and August Lemons vary much in flavour, the Messina lemons having the purest lemon flavour, those of Palermo having a slight additional musk and verbena odour, these and the Murcia lemons, which have a particularly fine flavour and are sometimes very large, weighing from 12 to 19½ oz, a large one yielding 2½ oz of peel, reach England from November to April A box of Messina Lemons contains 300 to 360 fruits, a box of Murcia Lemons only 200 Naples lemons are imported from April to August, but are inferior to the Sicilian kinds Malaga lemons, which have thick skins and numerous seeds, and proportionately less juice, arrive in September and October Lemons should be chosen with a thin rind and of medium size, as they are more juicy and are likely to keep better, the rind having been reduced in thickness by a process of curing or drying before packing, thus rendering them less likely to become mouldy

Tests—Lemon Peel yields not more than 5 p.c. of ash

Preparations.

OLEUM LIMONIS. OIL OF LEMON.

FR., ESSENCE DE CITRON, GER., CITRONENÖL; ITAL., ESSENZA DI CEDRO; SPAN., ESENCIA DE LIMON.

The Volatile Oil obtained from fresh Lemon Peel. A light yellow liquid with a pleasant odour and an aromatic, mild somewhat bitter after taste.

The Oil consists of about 90 per cent of the terpenes Dextro and Levo Limonene, the Dextro Limonene being the more important.

The terpene Phellandrene is also present in small quantity in the Oil. Pinene is not present. The odour and flavour of the Oil are dependent upon the oxygenated bodies, and are due chiefly to the aldehyde of Geraniol, Gerinaldehyde, $C_{10}H_{16}O$ eq 150.98, which is known commercially as Citral, and which is present to the extent of 3 to 4 per cent. See under Tests. A second aldehyde, Citronellal, is also present. An ester of Geraniol (Geranyl Acetate) is present in the Messina and Palermo Oils, whilst the latter Oil contains also a Linalyl ester (Linalyl Acetate) to which the difference in odour between the Palermo and Messina Oils has been partly attributed. A very small quantity of a sesquiterpene boiling at 240 to 242° C (464° to 467.6° F) is present. Octyl and Nonyl aldehydes have been detected in Lemon Oil and also Methyl Heptenone, Anthranilic Acid Methyl Ester, Citriplene and a Resin. Octyl and Nonyl aldehydes must be regarded as important constituents, as they play an important part in the aroma of the Oil. See also Citral.

Its flavour and aroma suffer much from keeping; it keeps the aroma much better if mixed (when fresh) with 10 per cent (by measure) of Absolute Alcohol. The presence of Ethylic Alcohol can readily be detected by the diminution in volume of the Oil on shaking with Water. The Oil should evaporate from paper without leaving a film.

Solubility. In all proportions of Glacial Acetic Acid and Absolute Alcohol, 1 in 12 of Alcohol (90 per cent).

Dose -- 1 to 3 minims = 0.3 to 0.18 cc.

Foreign Pharmacopœias. Official in Austri Oleum Citri, Belg, Essentia Citri, Dan, Aetherioleum Citri (Ger and Jap, Oleum Citri (sp gr 0.858 to 0.861), Dutch (sp gr 0.850 to 0.860), Hung (sp gr 0.840 to 0.870) Russ (sp gr 0.855 to 0.865), and Swiss (sp gr 0.857 to 0.861), all Oleum Citri, Ital, Essenza di Cedro? (sp gr 0.857 to 0.860), Fr, Essence de Citron (p gr 0.851 to 0.862), Mex, Accite Volatil de Limon (sp gr 0.849) Norw (sp gr 0.850 to 0.860) and Swed (sp gr 0.855 to 0.861), Aetherioleum Citri, Port, Essencia de Limao, (sp gr 0.846 to 0.856) Spain, Esencia de Limon (sp gr 0.852 to 0.856), U.S., Oleum Limonis (sp gr 0.851 to 0.855 at 25° C (77° F)).

Tests. Lemon Oil is officially required to possess a sp gr of 0.857 to 0.860, the *L.S.P.* requires 0.851 to 0.855 at 25° C (77° F), and the *P.G.* 0.858 to 0.861. The sp gr of the Oil is usually 0.856 to 0.858 and occasionally 0.860. The optical rotation is from +58° to +63 in a tube 100 mm in length. The official figure is not less than +59. The *L.S.P.* required that its optical rotation should not be less than +60 at a temperature of 25° C (77° F), this minimum rotation figure was considered too high, and with the 1907 season's Oil was virtually unattainable. The figure was altered in the list of Additions and Corrections (1907) to not less than +58°. The *P.G.* does not include a rotation figure. The refractive index of the Oil is not included in either the *B.P.*, *U.S.P.* or *P.G.*

It is from 1.473 to 1.483. Neither the *B.P* nor *P.G* requires the Oil to contain any definite percentage of Geranialdehyde (Citral), and no method of determination is given. The *U.S.P* requires that it shall yield not less than 4.0 p.c. of aldehyde calculated as Citral, when quantitatively determined by means of the following volumetric process, which is based upon the interaction of the aldehyde and a neutral 20 p.c. w/v Sodium Sulphite Solution, whereby Citral Dihydrosulphonic Acid is produced and a corresponding amount of Sodium Hydroxide is liberated, which is determined by titration with Semi-normal Volumetric Hydrochloric Acid Solution, using Rosolic Acid Solution as an indicator of neutrality. A control experiment with the reagents alone, without the Lemon Oil, is carried out simultaneously, the number of c.c. of Semi-normal Volumetric Acid required in this blank experiment being deducted from the number obtained in the actual determination. 1 c.c. of Semi-normal Volumetric Hydrochloric Acid Solution corresponds to 0.03802 gramme of Citral. Great diversity of opinion exists amongst authorities on essential oils, not only with regard to the actual amount of Citral contained in genuine Lemon Oils, but also with regard to the relative reliability of the various processes which have been suggested from time to time for its determination. The balance of opinion (more particularly amongst English authorities) appears to be that the standard of 5 to 7 p.c. of Citral cannot be maintained, and that the true percentage is nearer 3 to 4 p.c. Leaving chemical considerations out of the question altogether, Terpeneless oils are found to contain about 50 p.c. of Citral, which should mean that the yield of Terpeneless oil from Oil of Lemon should be about 14 p.c., whereas the manufacturers find that not more than half this amount is obtained. The determination of the aldehyde by the direct measurement of the portion unabsorbed by Sodium Bisulphite Solution is most conveniently carried out by mixing a measured quantity of 50 c.c. of the Oil with 75 c.c. of a 40 p.c. w/v Sodium Metabisulphite Solution and 25 c.c. of Sodium Sulphite (made by exactly neutralising the Sodium Bisulphite Solution with a 10 p.c. w/v Sodium Hydroxide Solution). The mixture is heated to 70° C (158° F) and shaken for one hour. Sufficient Water is then added to bring the Oil into the graduated portion of the flask and the unabsorbed portion is read off, a correction being made for the solubility of the Terpenes. This method has been adversely criticised, but is stated to give approximate results. The Hydroxylamine process has been found to give totally unreliable results. The Cyanacetic Acid method yields results which are invariably too high, and is useless when small percentages of Citral are concerned. The volumetric process with Sodium Bisulphite Solution (Sadtler's process), virtually that adopted by the *U.S.P*, is referred to above. Merck and Schimmel have adversely criticised the process, stating that the clearness of the end reaction is so poor as to be desired.

A method somewhat similar to Sadtler's process has been recommended (*C.D.* '05, n 408) for the determination of Citral. It is based on the reaction of Citral with Potassium or Sodium Sulphite

Solution containing sufficient excess of the Hydrogen Sulphate for the solution to remain acid after the absorption. The solution is prepared by dissolving 400 grammes of crystallised Potassium or Sodium Sulphite in 1 litre of Water and adding sufficient Potassium or Sodium Hydrogen Sulphite Solution to make each 25 c.c. sufficiently acid to neutralise 20 c.c. of Semi normal Potassium Hydroxide Solution. A measured quantity of 5 c.c. of the Oil is heated with sufficient of this solution in a closed flask for three hours on the water-bath and the residual acid is then titrated. Three molecular equivalents of acid are equivalent to one molecular equivalent of Citral, and consequently 1 c.c. of Semi normal Volumetric Potassium Hydroxide Solution corresponds to 0.02516 gramme of Citral. The process has been criticised by Messrs. Schimmel in their semi annual report, October to November, 1905, they find the process open to the same objection as Sadtler's process, viz., the indefinite nature of the end reaction, and did not succeed in the preliminary neutralisation of the solution as directed.

The more generally occurring adulterants of Lemon Oil are Turpentine Oil, Lemon Oil terpenes, Turpentine Oil to which Lemon Grass Oil Citral has been added, Turpentine Oils plus the addition of a little inferior Orange Oil, and Cedarwood Oil. Turpentine Oil is readily detected by distilling and examining the first 10 p.c. of distillate. This distillate is required to possess a rotation differing by not more than 2 from that yielded by the original Oil. This limit is considered very severe, genuine Lemon Oils often showing a greater difference. Lemon Oil terpenes are very difficult to detect. Their presence may be ascertained by fractional distillation and the determination of the optical rotation of fractions, and also by the reduction in the Citral content. To meet the reduction in Citral strength caused by the addition of Lemon Oil terpenes, Citral from Lemon Grass Oil has been added. It may be detected by the marked verbena odour communicated to the residue after about 90 p.c. of the Oil has been distilled off *in vacuo*. A method of fractional distillation, which it is claimed will show any adulteration, is given (J.S.C.I. '01, 1179). A measured quantity of 100 c.c. of the Oil is put into a distilling flask having three bulbs blown in the neck, and fitted with a cork and thermometer. It is connected with a condenser, fitted with a suitable receiver, having two vessels graduated at 10 c.c. and 80 c.c. respectively. It is exhausted, a pressure of not more than 15 mm. being maintained. The flask is gently heated by means of an oil-bath. The first 10 c.c. should not take more than seven minutes to distil. The next vessel is put into position, and the distillation continued until 80 c.c. have distilled over. The pressure is then relieved, and the residual oil in the flask is distilled over with steam, and the quantity obtained carefully noted. The optical rotation and the refractive indexes of the three fractions are determined, respectively, by the polariscope and Zeiss refractometer.

Turpentine Oil may be detected by the optical rotation, the diminution in the percentage of aldehydes, the effect upon the solubility of the Oil in Alcohol, and an examination of the portion

unabsorbed by Sodium Bisulphite. The presence of Citral from Lemon Grass Oil may be ascertained by a determination of the optical rotation and the marked verbena odour of the absorbed aldehydes.

The method adopted by Burgess and Child for ascertaining the solubility of a Terpeneless oil, and to which they refer as the solubility number is to dissolve a measured quantity of 1 cc of the Oil in 20 cc of Alcohol (94 p c), and then to add, from a burette, Distilled Water until a permanent milkiness ensues, the solubility number is the number of cc of Water required to produce such turbidity.

SYRUPUS LIMONIS SYRUP OF LEMON

Fresh Lemon Peel, in thin slices or grated, 1, Alcohol (90 p c), a sufficient quantity, Lemon Juice, 25, Refined Sugar, 38. Macerate the Lemon Peel in $1\frac{1}{2}$ of the Alcohol for seven days, press, filter, add sufficient of the Alcohol to produce 2. In the Lemon Juice, clarified by subsidence, dissolve the Refined Sugar by the aid of gentle heat. When the resulting syrup is cold, mix with it the 2 of Alcoholic liquid. The product should weigh 65.

(1 of Peel and 25 of Juice in 65)

This makes a turbid syrup. It is not possible to completely clarify it by subsidence. It has been suggested to filter the juice through Talc, but this removes not only the turbidity but also some of the flavour. It does not filter readily through flannel or paper, but if the juice is heated nearly to 212° F (100° C) and strained through flannel before dissolving the sugar, the resulting syrup is bright and clear.

Dose — $\frac{1}{2}$ to 1 fl drim = 18 to 36 cc

Foreign Pharmacopœias — Official in Fr (Syrup d'Acide Citrique) Citric Acid 1, Simple Syrup, 97, Alcoolature de citron, 2, Ital, Bruised Peel 2, Sugar 19, Distilled Lemon Water 12, Mex (Jarabe de Limon), Lemon Juice 10, Syrup 100, Port (Xarope de Casca de Limao), Fresh Lemon Peel 1, Boiling Water 35, Sugar 65, Span (Jarabe de Limon), Lemon Juice 5, Sugar 9, and Swiss, Citric Acid 2, Water 25, Syrup 94, Spirit of Lemon 15. For other Pharmacopœias, see Acidum Citricum.

TINCTURA LIMONIS TINCTURE OF LEMON

Macerate 5 of Fresh Lemon Peel with 20 of Alcohol (90 p c)

Now 1 in 4, B P 1885 was 1 in 8

Dose — $\frac{1}{2}$ to 1 fl drim = 18 to 36 cc

Foreign Pharmacopœias — in Belg (Spiritus Citri), 1 Oil in 100, Dutch (Spiritus Citri), 4 of Fresh Peel in 10 (distilled), Fr (Alcoolature de Citron), 1 Fresh Peel to 2 of Alcohol, Jap (Spiritus Citri), 1 Oil in 10, Mex (Alcoholato de Cortezas de Limon), Fresh Peel 2, Alcohol (80°) 10, Water 2, U S (Alcohol de Corteza de Limon), Peel 1, and Alcohol (80 p c) 6, distil, Swiss (Spiritus Citri), 12 of Fresh Peel in 100 (distilled), U S (Tinctura Limonis Cortex), Fresh Lemon Peel 1, Alcohol (95 p c), to produce 2. B P C (Tinctura Limonis Fortis), Fresh Lemon Peel 1, Alcohol (90 p c) 1.

Tests — Tincture of Lemon has a sp gr of 0.875 to 0.880, contains from 1 to 2 p c w/v of total solids and about 76 p c w/v of Absolute Alcohol.

Not Official

TERPENELESS OIL OF LEMON — A Terpeneless Oil of Lemon is an Oil from which practically the whole of the terpenes have been removed. When carefully prepared it is free from Limonene and contains very little of the Steroptene of the original Oil. The relative yield of the Terpeneless Oil is about 4 to 6 p.c. The Oil contains about 50 p.c. of Geranialdehyde (Citral), together with Citronellal, Geranyl Acetate, and Linalyl Acetate. Octyl and Nonyl Aldehydes, which are important constituents of Lemon Oil, require an additional importance in the Terpeneless Oil, on account of the modification they effect in the aroma and flavour of the Oil. Anthranic Acid Methyl Ester also forms a constituent of the Oil.

Tests — Terpeneless Lemon Oil has a sp. gr. of 0.895 to 0.899, an optical rotation in a 100 mm. tube of -5° to -8° , a refractive index in a Zeiss refractometer of 1.481 to 1.482.

The percentage of Geranialdehyde (Citral) should amount to not less than 40 p.c. nor more than 50 p.c. It may be determined by measuring the unabsorbed portion of the Oil after treatment with Sodium Bisulphite, or by the Sodium Sulphite method. Both processes are described in the large type under 'Oleum Limonis'.

The Oil may contain Terpenes due to imperfect separation or intentional adulteration, and Lemon Grass Oil Citral.

OILEM GRAMINIS CITRATI Oil of Lemon Grass. *Syn.* Indian Oil of Verbena. — The Oil distilled from *Andropogon Citratus*, dose $\frac{1}{2}$ to 3 minims = 0.08 to 0.18 c.c., is official in the *Ind* and *Col Add* for India, the Eastern Colonies and the West Indian Colonies.

CITRAL GERANIAL $C_{15}H_{24}O$, (q. 150.98) — A pale, yellow, mobile, optically inactive Oil, consisting of the high boiling point fractions from the distillation of Lemon Oil, having a penetrating lemon odour, and possessing a flavouring power about 15 times as great as the original Oil.

Sp. gr. 0.895 to 0.899, boiling point, 223° to 229° C.

It gives the aldehyde reactions with Bisulphites, and on reduction yields the alcohol Geraniol.

It may be used to increase the flavour of Oil of Lemon, by mixing it with the latter in the proportion of 1 to 14.

Teinture d'Essence de Citron Composée (Eau de Cologne) — Oil of Bergamot, 10, Oil of Orange, 10, Oil of Lemon, 10, Oil of Orange Flower, 2, Oil of Rosemary, 2, Alcohol (90 p.c.), 1000 — *Fr*.

LIMONIS SUCCUS.

LEMON JUICE

FR, SUC DE CITRON, *GER*, CITRONENSAFT, *IT*, SUCCO DI LIMONE,
SPAN, ZUMO DE LIMON.

The freshly expressed Juice of the ripe fruit of *Citrus Medica* var. *Limonum*. Contains 30 to 40 grams of Citric Acid to the fl. oz. It varies with the time of year. The acidity decreases as the season advances from November to April.

Lemon Juice is extremely liable to fermentation, and requires the addition of Alcohol to keep it, about 15 p.c. of Proof Spirit is sufficient.

Medicinal Properties — Refrigerant, when diluted, a particularly useful beverage in prevention and treatment of scurvy (3 or 4 oz. daily), relieves thirst in febrile and inflammatory affections. In acute rheumatism, $\frac{1}{2}$ to 1 pint = 284 to 568 c.c. daily.

Dose.—1 to 2 fl oz = 28 4 to 56 8 cc

Official Preparation—Syrupus Limonis, p 726 Used in the preparation of Acidum Citricum

Foreign Pharmacopœias—Official in Dutch (Succus Citri Artificialis), Citric Acid 1, Water 8, Spirit of Lemon 1, Fr, Mex (Jugo de Limones), Span (Zumo de Limon), US, from 7 to 9 pc of Citric Acid, Swiss (Succus Citri facticius), Citric Acid 10, Water 89, Spirit of Lemon 1

Tests—Lemon Juice has a sp gr of 1 030 to 1 040, contains from 10 to 14 pc w/v of total solids, and when evaporated to dryness and ignited leaves not more than 3 pc w/v of ash

It is officially required to contain from 7 to 9 pc w/v of Citric Acid, equivalent to from 30 to 40 grains per fl oz The acidity may be conveniently determined by titrating 10 cc of the juice with Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, about 10 1 cc will be required, 1 cc of Volumetric Sodium Hydroxide Solution = 0 0695 gramme of Citric Acid The USP requires that 10 cc of the juice shall neutralise at least 10 cc of Normal Volumetric Potassium Hydroxide Solution, using Phenolphthalein Solution as an indicator This corresponds to 6 95 pc of Citric Acid Lemon Juice is not official in the P G

No official tests are given for impurities It should evolve no odour of Sulphurous Acid when warmed When neutralised with Potassium Hydroxide Solution, then rendered faintly acid with diluted Hydrochloric Acid and shaken with Ether, the ethereal solution, when washed with a few cc of Water, should yield no purple-violet coloration when shaken with Water containing a drop of Ferric Chloride TS, showing the absence of Salicylic Acid Barium Chloride Solution added to the filtered juice should cause no turbidity or precipitate This test for Sulphuric Acid and Sulphates is included in the USP, as well as the following for Acetic and Tartaric Acids respectively When warmed with an equal volume of Sulphuric Acid and a few drops of Alcohol (94 9 pc) it should not evolve an odour of Ethyl Acetate (Acetic Ether) Upon the addition of a 1 in 3 Potassium Acetate Solution, and Alcohol (94 9 pc) it should not yield a white crystalline precipitate of Potassium Hydrogen Tartrate

100 cc of average Lemon Juice require for neutralisation about 11 4 grammes of Potassium Bicarbonate, about 9 5 grammes of Sodium Bicarbonate and about 16 5 grammes of Sodium Carbonate, or, if the Imperial quantities are adopted, 110 minims require for neutralisation about 11½ grains of Potassium Bicarbonate, about 9½ grains of Sodium Bicarbonate, and about 16½ grains of Sodium Carbonate

ACIDUM CITRICUM See ACIDUM CITRICUM

LINUM.

LINSEED

FR, LIN, GR, LLINSAMIN, ITAL, LINO, SPAN, LINO

The dried ripe Seeds of *Linum usitatissimum*, L.

The envelope or testa abounds in a peculiar gummy matter or mucilage, readily imparted to hot Water.

Medicinal Properties—Demulcent. Employed in faecal, pharyngeal and bronchial catarrh, dysentery, diarrhoea, and inflammatory affections of the urinary passages. In the form of Linseed Poultice it is applied to inflamed parts.

Official Preparations Linum Contusum and Oleum Lin.

Foreign Pharmacopœias—Official in Aust, Belg, Dutch, Fr. (Lin), Ger, Hung, Ital, Jap, Mex (Linaza), Norw, Port (Linlio), Russ, Span (Lino), Swed, Swiss and U.S.

Descriptive Notes Linseed varies much in size, the varieties imported from subtropical countries being distinctly larger than those cultivated in temperate or cold climates. Thus, of Russian, Dutch, English, and ordinary Calcutta Linseed, twelve or fourteen seeds, and of Archangel Linseed even seventeen seeds, weigh one gram, whilst of Bombay, Sicilian, and Ionian Linseed six or seven only are equal to a gram in weight, in other words, the last three are almost twice the size of the Linseed of temperate climates. For use in medicine the English and Dutch Linseeds are to be preferred, since they are usually most free from weed seeds, and from dirt. Most of the different varieties contain weed seeds distinctive of the country where they are produced, and can be recognised by these seeds, but any samples of Linseed containing more than 4 p.c. of weed seeds may be considered to be adulterated. Official Linseed is stated to be $\frac{1}{2}$ to $\frac{1}{4}$ in in length (4 to 6 mm), (3 to 5, USP), ovate and somewhat obliquely pointed, the surface glabrous and minutely pitted, of a brown colour, inodorous, with a mucilaginous taste. The mucilage exists in the epidermal cells, and is formed at the expense of starch, which is found only in the young cells of these cells. The seeds contain about one tenth of their weight of mucilage (6 p.c. Vogl), which can be precipitated in white flakes by Alcohol, but cannot be filtered until after boiling. Russian Linseed is largely used for the manufacture of Linseed Oil and for Linseed Cake, which is obtained by expression of the fixed Oil, and is employed for feeding purposes, since it contains up to 25 p.c. of proteids. Russian Linseed is often extensively adulterated with weed seeds, and even when sifted retains sufficient dirt attached to the surface to prevent the hand readily passing through a parcel of it, whereas English and Dutch permit it readily, and this test is used as a rough means of distinguishing these Linseeds. As the mucilage is contained in the epidermal cells, the seed is used in the whole state for making the decoction known as Linseed Tea. The principal features of powdered Linseed are the thin-walled, short prismatic cells, containing mucilage, yellowish spindle-shaped sclerenchymatous cells, crossed at right angles with a layer of

thin-walled, elongated, compressed, colourless cells, flattened polygonal cells with porous walls, containing a brown pigment, and endosperm cells, containing characteristic aleurone grains and drops of oil. It is best examined in strong Alcohol, and Water gradually added to show the mucilage cells.

Preparations

LINUM CONTUSUM CRUSHED LINSEED

Linseed reduced to a coarse powder

Foreign Pharmacopœias—Official in Belg, Fr and Ital, should contain 30 p c of Oil, —Official in U.S.P. and U.K., should contain 25 p c of Oil. Not in the others.

Tests—Crushed Linseed is officially required to yield, when extracted with Carbon Bisulphide, not less than 30 p c of Oil. This is the standard adopted by the *U.S.P.*, which also adds 'all of which is saponifiable'. The extracted Oil might with advantage be required to answer the tests for Oleum Lin. Good commercial samples, examined in the author's laboratory, yielded from 30 to 42 p c of oil. The *B.P.* statement that it should not yield the tests characteristic of Starch is very indefinite. Linseed contains a large proportion of Oil possessing a high Iodine absorption, if the Iodine test is applied, it is preferably carried out on the portion remaining after the extraction of the Oil with Carbon Bisulphide. The *U.S.P.* performs the test on the crushed Linseed, but gives very explicit instructions for carrying it out. A weighed quantity of 0.1 gramme is directed to be mixed with 20 c c of Water, the mixture heated to boiling, cooled, and diluted with cold Water to 100 c c. The addition of 0.5 c c of Iodine T.S. (2.0 p c w/v) should not produce more than a pale blue colour. The ash of crushed Linseed varies from 3 to 4 p c, and should not exceed 5 p c. It is officially required to leave, when incinerated with free access of air, not more than 5.0 p c of ash. No ash limit appears in the *U.S.P.* The *P.G.* includes the seeds but not the crushed seeds, but no tests are given. The 1890 *U.S.P.* required a yield of not less than 25 p c of Oil extractable by Carbon Bisulphide, which has been raised to 30 p c in the 8th Decennial Revision.

OLEUM LINI LINSEED OIL

A colourless, but more usually pale yellow, oily liquid, possessing a characteristic odour and unpleasant taste. It is a drying Oil, and tends to thicken and darken in colour on exposure to light and air. It is the Oil expressed from Linseed at ordinary temperatures.

For medicinal purposes it should be procured as fresh as possible.

Solubility—Of a freshly expressed sample, 1 in 40 of Absolute Alcohol, 1 in 1½ of Ether.

Medicinal Properties.—Laxative, it also acts mechanically as an enema for removing impacted faeces. A good application to burns in the form of Carron Oil, *see p. 289*.

Foreign Pharmacopœias—Official in Aust, sp gr 0.932 to 0.937, Belg, Ger, Hung, Norw, Russ, and Swed, sp gr 0.936 to 0.940, Dan, sp gr 0.930 to 0.940, Ind, sp gr 0.930 to 0.935, Ital, sp gr 0.935 to 0.940, Jap, sp gr 0.930 to 0.940, Port (Óleo de Linhaca), sp gr 0.930,

Span (*Acete de Linaza*), Swiss, sp gr 0.930 to 0.937 at 15° C and 0.880 to 0.881 at 98° C, U.S., sp gr 0.925 to 0.935 at 25° C (77° F) Hung, also *Oleum Lini Lotum*

Tests—Linseed Oil has a sp gr of 0.930 to 0.936, the *BP* states 0.930 to 0.940, the *USP* 0.925 to 0.935 at 25° C (77° F), *PG* 0.936 to 0.940. All three Pharmacopœias are agreed that it does not congeal at temperatures above -20° C (-4° F). It is a drying Oil, and gradually thickens by exposure to the air, forming a hard transparent varnish, this feature is recorded in the *BP* and *USP*, but not in the *PG*. Neither the Saponification value nor the Iodine absorption is given in the *BP*, both should be included. The Saponification value usually ranges from 190 to 195, and should not be less than 187, and the Iodine absorption from 170 to 188. The *USP* gives the Saponification value as 187 to 195 and the Iodine value as not less than 170. The *PG* does not include a Saponification value, but requires that the Oil shall absorb not less than 150 p.c. of Iodine.

The more generally occurring adulterations are Mineral Oils, Rosin, and Rosin Oils. Mineral and Rosin Oils are unsaponifiable, and their presence may be detected by the Potassium Hydroxide test described below. If after saponification the Alcohol be evaporated off and the residual soap be dissolved in hot Water, cooled and shaken with Ether, the ethereal solution may be separated, evaporated to dryness and the unsaponifiable residue weighed. Rosin Oil may be detected by dissolving a portion of this residue in Carbon Bisulphide and adding a few drops of a Carbon Bisulphide Solution of Stannous Bromide containing an excess of Bromine. The appearance of a violet or purple coloration is indicative of the presence of Rosin Oil. The reagent may be prepared by allowing Bromine to fall drop by drop on some granulated Tin until the permanent coloration of the product shows that the Bromine is in excess. A further moderate quantity of Bromine is then added, and the mixture, when cold, diluted with three or four times its measure of Carbon Bisulphide.

The *USP* includes a test with Glacial Acetic Acid (see below) for the detection of Rosin Oils. The Acid should not be more than faintly acid in reaction towards *Ph* litmus paper moistened with Alcohol (94.9 p.c.). The Acid value should be less than 5, though old oil may give figures as high as 7. Linseed Oil contains a small percentage of unsaponifiable matter, which should not amount to more than 2.5 p.c., and usually is considerably less. Linseed Oil, when issuing from the seed without pressing, has scarcely any of the odour or taste of the Linseed Oil of the shops, but acquires it in a very short time on exposure to the air. For medicinal purposes it should be procured as fresh as possible. Boiled Linseed Oil is used in the arts as a drying Oil, and for certain purposes Litharge and Manganese are added during the boiling. The boiled Oil may therefore contain both Lead and Manganese. It may be detected by the increased sp gr and the great decrease in the Iodine absorption.

The rise of temperature on treating the Oil with Sulphuric

Acid (Maumené's test) is a useful constant for the Oil, it should not be less than 114°C (237°F)

Potassium Hydroxide—If 27 parts of Potassium Hydroxide T S and 2 parts of Alcohol (90 p c) be added to 20 parts of Linseed Oil, warmed in a deep tin or porcelain vessel, and the mixture stirred and again warmed until a saponification is complete, the soap formed should be soluble in Water or Alcohol (90 p c) without residue, *P G*, it should be completely saponifiable with Alcoholic Potassium Hydroxide T S, and the resulting soap should be completely soluble in Water without leaving an oily residue, *U S P*

Glacial Acetic Acid—2 c c of the Oil warmed and shaken in a test tube with an equal volume of Glacial Acetic Acid should yield, on cooling and the addition of 1 drop of Sulphuric Acid, a greenish colour. A violet colour indicates the presence of Rosin or Rosin Oils, *U S P*

Not Official

CATAPLASMA LINI—Linseed Meal, 4, Boiling Water, 10. Mix the Linseed Meal with the Water gradually, with constant stirring. In cold weather the basin should be previously rinsed with boiling Water.
Applied to inflamed parts

Foreign Pharmacopœias—Official in Fr, Mex, Port and Span

Not Official.

LITHIUM

Li, eq 6.97

A silver-white, brilliant, ductile metal, having the density of 0.59

It is obtained from several minerals—Petalite, Lepidolite, Triphane, and formerly from Triphylline

Official Preparations—Lithii Carbonas, Lithii Citras

Not Official—Lithii Berizoas, Lithii Bromidum, Lithii Guaiacas, Lithii Hippuras, Lithii Quinas, Lithii Salicylas, Lithii Theobrominæ Salicylas, Lithii Bitartaras

Tests—Lithium has a sp gr of 0.59. The characteristic test for Lithium is the crimson colour which its salts, especially when moistened with Hydrochloric Acid, communicate to a non-luminous flame. An aqueous solution of a Lithium salt affords a precipitate with Sodium Phosphate Solution

LITHII CARBONAS.

LITHIUM CARBONATE

Li_2CO_3 , eq 73.49

FR, CARBONATE DE LITHIUM, GER, LITHIUMCARBONAT, ITAL, CARBONATO DI LITIO, SPAN, CARBONATO DE LITINA O LITICO.

A light white, amorphous, odourless powder, alkaline in reaction. It may be obtained from native Lithium Silicates

The salts official in the *BP* and the *USP* are required to contain not less than 98.5 p c of pure Lithium Carbonate, the *P. G.* does not specify the percentage

It should be kept in well-stoppered glass bottles.

Solubility—About 1 in 70 at 60°F . In hot Water it is only soluble to about half this extent, a solution saturated in the cold

becoming quite turbid on boiling. It should be noticed that using 1 part of Lithium Carbonate to 70 parts of Water solution is very slow, and using these proportions in oz it requires several weeks' digestion, with frequent shaking, before complete solution is effected.

Medicinal Properties—Diuretic. Combined with Carbonic Acid, in a diluted solution, as in Lithia Water, it has been given in cases of gout with the view of increasing the alkalinity of the blood, and acting as a solvent of the Sodium Bicarbonate deposits.

Luff has shown that the Lithium salts do not exercise any special solvent effect on Sodium Bicarbonate, and that their administration to gouty subjects with the object of removing uratic deposits in the joints and tissues appears to be useless—*L* '98, i 1609.

He also found that Lithium salts, although they did not delay the initial conversion of the gelatinous Sodium Bicarbonate into the crystalline forms, yet when the conversion was once started it was slowed by the presence of these salts and especially by the Lithium Carbonate. In the treatment of gout the Potassium salts were the most useful, and the Lithium salts ranked next—*L* '00, i 981, *J. M. J.* '00, i 536.

Cases of cardiac depression and even dilatation, as the result of the excessive and continued consumption of Lithia tablets, which are so persistently vaunted as curative of gout—*Ph* '07, i 166.

1 grain of Lithium Carbonate with $\frac{1}{2}$ grain Sodium Arsenate given in aerated Water has been recommended by Martineau, in the treatment of diabetes—*L* '87, i 650.

Dose—2 to 5 grains = 0.13 to 0.32 gramme.

Prescribing Notes—Given in aerated Water, cachets, or Compressed Tablets. *Varalettes* are effervescent tablets.

For the granular effervescent form, see *Lithia Citras*.

Official Preparation—Used in the preparation of *Lithia Citras*.

Not Official—Liquor *Lithii Carbonatis*.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S. Span., also *Carbonato de Litina efervescente*.

Tests—Lithium Carbonate dissolves slowly in Water and yields a solution which has an alkaline reaction towards red Litmus paper. It dissolves in diluted Hydrochloric Acid with effervescence, yielding a solution which gives the tests distinctive of Lithium given under that heading, and a colourless gas which, passed into Lime Water, affords a white precipitate, which again dissolves if a sufficient excess of the gas be passed through the liquid, or which is soluble with effervescence in Hydrochloric Acid. It is officially required to yield 98.5 p.c. of Lithium Carbonate as gravimetrically determined by neutralising 1 gramme of the salt with Sulphuric Acid, and subsequently heating to redness. The weight of dried Lithium Sulphate should amount to 1.479 grammes. It should be noted that the weight of Lithium Carbonate corresponding to 1.479 grammes of Lithium Sulphate is 0.9946 gramme, calculating out to 99.46 p.c. and not, as officially stated, 98.5 p.c. The *U.S.P.* and the *P.G.* employ volumetric methods of determination. The *U.S.P.* requires the salt to yield 98.5 p.c. of pure Lithium Carbonate as volumetrically determined by the process given in the small type below. The *P.G.* process, also given below, is direct titration. It is performed on the

salt dried at 100° C (212° F) and indicates 99.24 p.c. of Lithium Carbonate

The more generally occurring impurities are Aluminium, Ammonium, Arsenic, Copper, Iron, Lead, Magnesium, Potassium, Sodium, Zinc, and Chlorides, for which it is officially required to yield no characteristic reaction, and Calcium and Sulphates, for which it is required to yield only the slightest reactions. The salt when dissolved in diluted Acetic Acid should leave no insoluble residue. In carrying out the following tests the salt may be dissolved in diluted Hydrochloric Acid. For the detection of Iron and Aluminium may be detected by the addition of Ammonia Solution as described in the Ammonia test below. Arsenic, Copper, and Lead may be detected by the Hydrogen Sulphide test, Zinc by the subsequent addition of Ammonia. A standard for Lead of 10 parts per million is suggested (C.P. '08, p. 796), and 2 parts per million for Arsenic. Magnesium may be detected by the usual group reagents, Ammonium, by the behaviour on boiling with Liquor Potassæ, Potassium and Sodium in the residue after separation of all other metals and evaporation to dryness. Calcium, Chlorides and Sulphates, may be detected by the tests given below in small type, with Ammonium Oxalate, Silver Nitrate, and Barium Nitrate Solutions, respectively. The P.G. requires that 0.2 gramme of Lithium Carbonate dissolved in 1 c.c. of Hydrochloric Acid and evaporated to dryness, should leave a residue which yields a clear solution in 3 c.c. of Alcohol. The U.S.P. uses the Amyl Alcohol test, as described below, in fixing a limit of other alkalis.

Acetic Acid—A solution (1:50) of the salt dissolved in 40 c.c. of diluted Acetic Acid should leave no insoluble residue. U.S.P.

Ammonia—1 part of Lithium Carbonate mixed with 20 parts of Water and Hydrochloric Acid added drop by drop until the salt is dissolved yields a solution which, on the addition of T.S. of Ammonia until it is of alkaline reaction, should produce no turbidity nor precipitation either before or after boiling, U.S.P.

Hydrogen Sulphide—An aqueous solution of the salt (1:50) obtained by the aid of Nitric Acid after the addition of excess of T.S. of Ammonia, should be unaffected by T.S. of Hydrogen Sulphide, P.G., a solution obtained as directed in the preceding U.S.P. test, should not respond to the time-limit test for heavy metals, U.S.P.

Silver Nitrate—A solution (1:50) of the salt as above should not become turbid more than opalescent with T.S. of Silver Nitrate, P.G.

Barium Nitrate—A solution (1:50) of the salt as above is unaffected by T.S. of Barium Nitrate, P.G.

Ammonium Oxalate—A solution (1:50) of the salt as above is unaffected by T.S. of Ammonium Oxalate, P.G.

Amyl Alcohol—If 0.2 gramme of the salt contained in a flask of 50 c.c. capacity a slight excess of Hydrochloric Acid be added and the mixture evaporated almost to dryness on a water bath, and if 10 c.c. of Amyl Alcohol [boiling point 132° C (269° F)] be added and the mixture cautiously heated, the layer has evaporated, then upon the addition of 3 drops of Amyl Alcohol and boiling for three minutes the resulting insoluble residue should weigh not more than 0.003 gramme. The removal of the Water from the Amyl Alcohol is facilitated by passing a current of air through the hot solution. U.S.P.

Volumetric Determination—0.5 gramme of Lithium Carbonate dried at 100° C (212° F) should require for neutralisation not less than 13.4 c.c. of

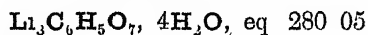
Normal Volumetric Solution of Hydrochloric Acid, *PG*, a solution of 0.5 gramme of Lithium Carbonate in 20 c.c. of Normal Sulphuric Acid Volumetric Solution should require not more than 6.6 c.c. of Normal Potassium Hydroxide Volumetric Solution for complete neutralisation, using Methyl Orange Solution as indicator, *USP*

Not Official

LIQUOR LITHII CARBONATIS (Lithia Water) —10 fl. oz. of aerated Water contain 5 grains of Lithium Carbonate

LITHII CITRAS.

LITHIUM CITRATE



A white crystalline powder, possessing a cool, slightly alkaline taste

It may be prepared by neutralising Citric Acid with Lithium Carbonate

It has been recommended that the formula should be altered to $\text{Li}_3\text{C}_6\text{H}_5\text{O}_7, 5\text{H}_2\text{O}$, eq. 297.93. The 8th Decennial Revision of the *USP* maintains the same formula as the *BP* 1898. Dott states (*CD* '05, 1 489) that the formula with four molecules of Water of crystallisation is generally accepted as accurate. It has been pointed out (*CD* '05, 1 489) that the *BP* is inaccurate in describing the salt as deliquescent. *USP* describes it as deliquescent in moist air.

Solubility —1 in 2 of Water, almost insoluble in Alcohol (90 p.c.)

The solubility in Water is variously given as 1 in 5 to 1 in 25.

Medicinal Properties —Similar to those of the Carbonate, but the Citrate being more soluble, it is better adapted for fluid administration.

Dose —5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes —Generally given in solution, or in the form of *Lithii Citras Effervescens*

Official Preparation —*Lithii Citras Effervescens*

Not Official —*Lithii Citras Lavativus Effervescens*

Foreign Pharmacopœias —Official in U.S., Mex. (*Citrato de Litio*) Not in the others

Tests —Lithium Citrate responds to the tests distinctive of Lithium given under that heading. A 5 p.c. aqueous solution of the salt yields, when boiled with an excess of Calcium Chloride Solution, a white precipitate insoluble in Potassium or Sodium Hydroxide Solution, soluble in Ammonium Chloride Solution. Its aqueous solution is generally faintly alkaline in reaction towards red Litmus paper, but should not redden Phenolphthalein Solution.

It is officially required to lose 19.0 p.c. of moisture when dried at 100° C (212° F) and an additional 6.5 p.c. at a temperature of 115.5° C (240° F). This statement has been shown (*CD* '05, 1 489) to be incorrect. Well-defined crystals, apparently quite dry, lose

considerably more than 19.0 p.c. at 100° C (212° F). Some large crystals, after exposure for two days to the air, lost 24.8 p.c. under 100° C (212° F), and when heated to a temperature of 150° to 160° C (302° to 320° F) lost a further 2.8 p.c.

The *B.P.* requires 2 grammes of the salt to leave when burned at a low red heat with a free access of air 0.77 gramme of white residue, corresponding to 98.5 p.c. of the pure Citrate. It is an extremely difficult matter to obtain a white residue, when burned at a low red heat, and, assuming the residue to consist of Lithium Carbonate, the *Pharmacopoeia* requires 97.80 p.c. of pure Lithium Citrate, and not less than 97.5 p.c. The *U.S.P.* gravimetric method of determination requires that the salt shall contain not less than 98.44 p.c. nor more than 100.2 p.c. of pure Lithium Citrate. The Carbonate left on cautious ignition is converted into Sulphate, and the complete oxidation of the carbonaceous residue ensured by cautiously re-igniting the residue with a few drops each of Nitric and Sulphuric Acid. The Citrate is not official in the *P.G.*

The Citrate being prepared from Lithium Carbonate, the impurities present in the latter are also liable to be present in the former, and the same methods as are adopted for their detection may also be employed, *see* Lithium Carbonate. Standards are suggested (*C.D.* '08, 1796) of 5 parts per million for Lead, and of 1 part per million for Arsenic. The tests for Iron and Aluminium may be carried out on the residue left on ignition after neutralisation with Hydrochloric Acid.

Gravimetric Determination—0.5 gramme of Lithium Citrate, dried at 150° C (302° F) cautiously ignited in a porcelain crucible, the residue cooled, then moistened with a few drops of Nitric and Sulphuric Acids and again cautiously ignited, repeating this operation until the residue of Lithium Sulphate becomes white and of constant weight, it should weigh not less than 0.387 gramme nor more than 0.394 gramme, *U.S.P.*

Ammonia—Dissolve the residue of Lithium Sulphate from the gravimetric determination in 10 c.c. of boiling Water and add Hydrochloric Acid. The addition of Ammonia Water until the solution has an alkaline reaction should not cause a turbidity or produce a precipitate either before or after boiling, *U.S.P.*

Time-limit Test—The (1-20) solution acidulated with Hydrochloric Acid should not respond to the time-limit test for heavy metals, *U.S.P.*

Amyl Alcohol—Let the residue obtained by the gravimetric test, the salt at a red heat be treated with a slight excess of Amyl Alcohol, and the mixture filtered. Then if the filtrate and washings be evaporated and further treated as described under the same heading under Lithium Carbonate, the resulting insoluble residue should weigh not more than 0.002 gramme, *U.S.P.*

Preparation

LITHII CITRAS EFFERVESCENS EFFERVESCENT LITHIUM CITRATE

Sodium Bicarbonate, in powder, 58, Tartaric Acid, in powder, 31, Citric Acid, in powder, 21, Lithium Citrate, 5 make into granules (1 in 20)

Dose—60 to 120 grains = 4 to 8 grammes.

Foreign Pharmacopœias—Official in U.S. Not in the others.

Not Official

LITHII CITRAS LAXATIVUS EFFERVESCENS—Lithium Citrate, 10, Sodium Phosphate, dried, 30 Sodium Bicarbonate, 44, Tartaric Acid, 15, Citric Acid, 17 50—*Bournemouth Formulary*, and *B P C*

LITHII BENZOAS ($\text{Li C}_7\text{H}_5\text{O}_2$, eq 137 10)—A white powder, or small shining scales, with a faintly acid reaction, the taste is sweet and somewhat saline. It can be prepared by boiling, in Water, 73 49 parts of Lithium Carbonate with 242 26 parts of Benzoic Acid, and evaporating.

It should contain not less than 98 5 p.c. of pure Lithium Benzoate.

It should be kept in well stoppered bottles and in a cool atmosphere.

Solubility—1 in 24 of Water, 1 in 15 of Alcohol (90 p.c.)

The solubility of the salt varies with the amount of uncombined Benzoic Acid which it contains. A pure salt prepared by exactly neutralising Lithium Carbonate with its ascertained equivalent of Benzoic Acid gave the above figure. The figure 1 in 14 recorded in the *Pocket Companion* was obtained from a sample containing 8 6 p.c. of free Benzoic Acid. See also under Tests.

Medicinal Properties—Used extensively in gouty conditions.

Dose—15 to 30 grains = 1 to 2 grammes.

Foreign Pharmacopœias—Official in *Fr*, *Ital*, *Span* (*Benzoato Litico*), and *U S*, *Mex* (*Benzoato de Litio*). *Span* has also *Benzoato Litico effervescente*. Not in the others.

Tests—Lithium Benzoate fuses when heated, and at a higher temperature chars, evolving vapours having a Benzoin odour, and finally burns away leaving a residue of Lithium Carbonate. It responds to the tests distinctive of Lithium given under that substance. Its aqueous solution is alkaline in reaction towards red Litmus paper, and, if well prepared, is neutral in reaction towards Phenolphthalein Solution, but commercial samples are frequently acid in reaction towards Phenolphthalein Solution, requiring an appreciable quantity of Tenth normal Volumetric Potassium or Sodium Hydroxide Solution to restore neutrality.

Samples examined in the author's laboratory have contained from 6 to 14 p.c. of uncombined Benzoic Acid. A 5 p.c. aqueous solution yields with Ferric Chloride T.S. a buff coloured precipitate. A concentrated aqueous solution affords with Hydrochloric Acid a white precipitate soluble in Ether and in Potassium and Sodium Hydroxide Solution. If this precipitate be separated, washed till free from mineral acid, and carefully dried it should possess the m.p., respond to the tests for and be free from the impurities mentioned under 'Acidum Benzoicum'.

The salt is official in the *U S P* and is required to contain not less than 97 66 p.c. nor more than 100 2 p.c. of pure Lithium Benzoate as gravimetrically determined by cautiously igniting in a porcelain crucible a weighed quantity of the salt with about twice its weight of powdered anhydrous Ammonium Sulphate, the weight being recorded when constant. The *U S P* 'purity rubric' says it shall contain not less than 98 5 p.c. of pure Lithium Benzoate, which does not correspond with the above gravimetric determination. The percentage of Lithium Benzoate may also be determined from the alkalinity of the residue on ignition, and this method was adopted in *U S P* 1300. The process was not accurate, and was tedious, on account of the difficulty of burning off the carbonaceous matter. The author has found the following method of direct titration both expeditious and accurate.—Dissolve a weighed quantity of 1 gramme of the salt in 50 c.c. of pure Distilled Water, add about 10 c.c. of Ether and a few drops of Phenolphthalein Solution, shake and titrate with Tenth normal Volumetric Sodium Hydroxide Solution, 1 c.c. of the Tenth normal Alkali Solution = 0 012113 gramme Benzoic Acid. A few drops of Methyl Orange Solution are added and the titration continued with Tenth normal Volumetric Sulphuric Acid Solution, 1 c.c. of which = 0 01271 gramme Lithium Benzoate.

LITHII BROMIDUM (Li Br , eq 86 32)—A white, granular, deliquescent salt, having a sharp and somewhat bitter, saline taste.

It should contain not less than 97 p.c. of pure Lithium Bromide.

It should be kept in well-stoppered glass bottles of a dark amber tint and in a cool place

Solubility—1 in 1 of Water, 1 in 4 of Alcohol (90 p c)

Medicinal P the low atomic weight of Lithium, this salt contains no Potassium or Sodium Bromide, and consequently has been recommended as a hypnotic for gouty patients, and in epilepsy

In the insomnia of 30 grains three times a day) P, 1: 351
In Bright's disease—L n gouty cases of aural vertigo, especially when preceded by a mercurial purge—M A '95, 221

Dose—5 to 15 grains = 0.32 to 1 gramme

Foreign Pharmacopœias—Official in Russ and U S, Mex, Bromide de Litio Not in the others

Tests—Lithium Bromide fuses when heated at a low red heat, and is slowly volatilised at a higher temperature. It responds to the tests distinctive of Lithium given under that heading. It is slightly alkaline in reaction towards red Litmus paper, and with Nitric Acid affords with Silver Nitrate Solution a yellowish curdy precipitate, insoluble in Nitric Acid, practically insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution. Chlorine Water added cautiously to the aqueous solution affords a yellowish coloration, and if the liquid be shaken with Chloroform the colour passes into the Chloroform solution. The salt is official in the U S P and is required to contain not less than 97.13 p c nor more than 101.17 p c of pure Lithium Bromide as volumetrically determined by dissolving a weighed quantity of the dried salt in a measured quantity of Water and titrating an aliquot portion of the solution with Tenth-normal Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator.

The more generally occurring impurities are those also mentioned under Lithium Carbonate, and similar tests may be employed for their detection. The U S P includes a test for Potassium with Sodium Cobaltic Nitrite Solution, prepared by dissolving 4 grammes of Cobaltous Nitrate and 10 grammes of Sodium Nitrite in about 50 c c of Water, adding 2 c c of Acetic Acid and diluting with sufficient Water to produce 100 c c, 0.5 c c of this solution added to 5 c c of a 5 p c aqueous Lithium Bromide Solution should not in 10 minutes produce either a turbidity or precipitate. Iodine may be detected by a violet coloration imparted to the chloroformic or Carbon Bisulphide Solution of the Bromine liberated when the aqueous solution of the salt is treated with Chlorine Water and shaken with either of these solvents.

LITHII GUAÏACAS.—Is prepared by Guaiacum Resin in an aqueous solution of Lithium Oxide, decanting solution, evaporating and scaling it. Composed of Lithium Oxide, 1, Guaiacum Resin, 3
(Given for chronic gout and some forms of rheumatism)

Tests—Lithium Guaiacate responds to the tests distinctive of Lithium given under that heading. The aqueous solution yields a blue coloration on the addition of a drop of Ferric Chloride T S.

Dose—5 grains = 0.32 gramme, twice a day, made into a pill with Dispensing Syrup.

LITHII HIPPIURAS—A white micro-crystalline powder, soluble 1 in 2½ of Water. It has been used as a solvent for Uric Acid deposits.

Dose—5 to 15 grains = 0.32 to 1 gramme

Tests—Lithium Hippurate yields the tests distinctive of Lithium given under that heading. Its neutral aqueous solution yields on the addition of Hydrochloric Acid an immediate crystalline deposit, and with Ferric Chloride T S a cream coloured precipitate, soluble in excess of the reagent.

LITHII QUINAS (Lithium Kinate Urosino)—A whitish or brownish white, granular sum of tablets. A 50 p c solution is also supplied for a solvent of Uric Acid deposits in gout—B V J '91, 1: 170; G L, 1: 478; I, '99, 1: 1722; P J '00, 1: 57.

combined with other purgatives
tive for children

Prescribing Notes—It frequently
solid mass when prescribed in mixtures,
Magnesium Carbonate is not open to the

Although the heavy powder is preferred
powder is said to be quicker in its action

It may be given in Water or in Milk
added at once, so as to form a smooth paste

Dose—5 to 30 grains = 0.3 to 2 grammes
stration, for a single administration

Incompatibles—All acids

Official Preparation—Permitted

Foreign Pharmacopœias—when strongly required to contain
Ponderosa), Norw and Swed (Oxid unburnt and unburnt)
US Not in the others

Tests—Heavy Magnesium Sulphuric Acid forming a solution of Magnesium given under the name of Light Magnesium Oxide by a precipitate when 1 part of the sample should otherwise conform to the test. It is liable to contain the same methods may be used as are described under the name of Magnesium Sulphate.

MAGNESII

LIGHT MAGNESII

3(MgCO₃),

FR, HYDROCARBONATE OF MAGNESIUM, CARBONATO DI MAGNESIO

A very light, white, odourless substance of Iron salts, (see USP and PG incl.)

It has been stated that within certain limits and conditions The Light Carbonate alone have a formula 'approximate' 482.26 It is required

of residue, of which none of Magnesium Oxide 1 oz occupies about

Solubility—1 in 20 of Water

Medicinal Properties—0.32 to 1 gramme

Prescribing Notes—0.32 to 1 gramme

Dose—5 to 30 grains = 0.32 to 2 grammes
stration, for a single administration

It should be kept in well-stoppered glass a cool place

Solubility—1 in 1 of Water, 1 in 4 of A-ture of 0.2 grammes of

Medicinal Properties—Owing to the less than opalescent within 5 minutes with salt contains more Bromide than either Pot consequently has been recommended as a hy

ution of 1 gramme of Manganese in 100 c.c. of Water should be colourless

In the insomnia of the aged (30 grains) with T.S. of Potass. In Bright's disease—*L* 1, 2, 3. In gouty 20) prepared with when preceded by a mercurial purge!—*M A* 95) tion with T.S. of Potassium Ferro-

Dose—5 to 15 grains = 0.32 to 1 gramme

Foreign P. and connections Official in P4 in the Vignesia in 10 cc of
de Lizio Not T-0 H-1000 S-1000 P-6 A

Tests—Lithium Bromide fuses when heated, should not respond to flame test, volatilised at a higher temperature. It responds to the test for bromine.

Lithium given under that heading. Its aqueous Acetic Acid as above should not
reaction towards red Litmus paper, and when acetic Nitrate, P/G

with Silver Nitrate Solution a yellowish curdy, and not, after the addition of Nitric Acid, insoluble in Ammonia Solution with Silver Nitrate TS within 5

Cyanide Chlorine Water added cautiously affords a yellowish coloration, and if the liquid colour passes into the Chloroform solution. Th⁰ grammes of recently ignited and

and is required to contain not less than 97.13 per cent Volumetric Sulphuric Acid
pure Lithium Bromide as volumetrically determined using Methyl Orange TS as

quantity of the dried salt in a measured quantity, using Methyl Orange TS as an equal portion of the solution with Tenth-normal Sulfuric Acid Solution is equivalent to Solution, using Potassium Chromate Solution as a

The more generally occurring impurities are with 15 parts of Water in a beaker Lithium Carbonate, and similar tests may be employed. If it will form a gelatinous mass, it is Lithium Carbonate. For Potassium, with Sodium, but when the beaker is

USP for Potassium with Sod (p. 11), out when the beaker is prepared - 4 grammes of Cobaltous with Water to form a gelatinous Sodium Nitrite in about 50 cc of Water, add

diluting with sufficient Water to produce 100 c c ,
to 5 c c of a 5 p c aqueous Lithium Bromide Solution
make a white turbidity or precipitate. Induce

produce either a turbidity or precipitate. Iodine
coloration imparted to the chloroformic oil. Car
Bromine liberated when the aqueous solution of **EROSA.**

EROSA.

LITHII GUAIACAS—Is prepared by digesting aqueous solution of Lithium Oxide, decanting the clear liquid, and evaporating to dryness. Composed of Lithium Oxide 1. Guaiacum 2. Glycerine 1. Magnesia 1.

Tests—Lithium Guaiacate responds to the tests for lithium and having an earthy and

given under that heading. The aqueous solution yielded having an earthy and addition of a drop of Ferric Chloride T.S. heavy M. green in Car.

LITHI HIPPIURAS—A white micro-crystalline salt—closed glass, by con-

Dose.—5 to 15 grains, = 0.32 to 1 gramme

Tests.—Lithium Hippurate yields the tests distinguished from in hot Water under that heading. Aqueous solution yields the tests for Lithium. Much more soluble in water than in alcohol.

LITHIUM SALTS (Lithium Salts) — (ended with acid) and

white, granular, effervescent powder, or in the form of tablets, it may often be used as a solvent. Employed as a solvent, it is not so effective as the other agents mentioned. It is not so effective as the other agents mentioned. It is not so effective as the other agents mentioned.

Foreign Pharmacopœias—Official in Ger, Hung, Ital, Jap, Mex, Norw, Pol, Fr has also Hydroxide de Magnesium

Tests—Light Magnesia when affords a solution which answers given under that heading. Accordingly distinguished from Heavy Magnesia Hydroxide when 1 part of the Water. It is not officially required of pure Magnesium Oxide. The ignited and cooled Oxide shall contain Magnesium Oxide, as volumetrically indicated below. The *PG* gives

The more generally occurring excess of moisture, Aluminium, and Sulphates. The presence of by the test with Litmus and type below. Excess of moisture sample is heated to a dull red lose little or no weight, the Water of hydration. Aluminium of a precipitate on the neutralised solution of the taining an excess of Ammonium a separate test for Iron. Calcium is shown by the Chlorides, and Sulphates. Silver Nitrate tests respectively Arsenic, Copper, Iron, Lead adopting the time limit test and Lead from the official Magnesium Carbonate is remarkable, highly objectionable been found.

Standards of 4 parts million for Lead have been is mentioned that many especially is thus true. **OMINÆ SALICYLAS** instance in the paper suggested.

Litmus—If 1 grammes = 1 to 1.3 grammes to boiling, cooled and filtered. than a faintly alkaline reaction. 0.2 grammes of Magnesium yields a filtrate at most but.

Residue—The filtrate, RAS (Lithium Acid Tartrate) rated to dryness should not be 5 cc of the filtrate obtained leave only a very insignificant

Acetic Acid—0.1 grm Citrate and Sodium Sulphate of Water, then cooled and pleasant laxative action than tion without evolution of rains = 4 to 8 grammes

It should be kept in well-stoppered bottles, added 100 grains of Magnesium Carbonate, stir until it is a cool place solution into a strong half-pint bottle, add $\frac{1}{2}$ fl oz of Syrup

Solubility—1 in 1 of Warm Bicarbonate in crystals, and immediately close the bottle

Medicinal Properties—should be secured with stung or wine, afterwards shake the salt contains more Bromide, Cassium Bicarbonate is dissolved consequently has been recorded as aperient and refrigerant draught
0 fl oz = 142 to 284 c c

In the insomnia of neu Carbonate, 15, Citric Acid, 33, Syrup of Citric Acid,* 60, In Bright's disease—L⁹⁵ bonate, 2 5, Water, q s to make about 360 — U S P when preceded by a mercuri incorporated in the B P C

Dose—5 to 15 grains citri Citrici—Citric Acid, 1, Distilled Water, 1, Tincture of Foreign Pharmacopoeias, 1, Syrup, q s to produce 100 — U S P

de Latio Not in the other Pharmacopoeias—Official in the U S formula modified Austr

Tests—Lithium Brotio Magnesiæ Citricæ Effervescentis, Austr has also volatilised at a highern Citricum Effervescentis, Belg (Magnesiæ Citrici Lithium given under Dutch (Solutio Citratis Magnesiæ), Fr (Limonade reaction towards red Lignéssionne), Ital (Limonata Magnesiaca), Mex (Solu- with Silver Nitrate Solutio de Magnesia), Port (Limonada Citrio-Mag- Acid, practically insoluble (Potion Magnesiæ Citricæ Aerophoria), Spm (Potion Cyanide Solution Co Magnesiæ Gaseosa), also Poción de Citrato de Mag- affords a yellowish col, Jap and Swiss (Magnesium Citricum Effervescentis) colour passes into the also Limonata aerata laxans Not in the others Fr has and is required to cate de Magnesie desseche

pure Lithium Bion **MAGNESIUM LACTATE**—Valuable for combating the accidents of quantity of the dr has in cases where the Calcium salts do not seem to act Dose, 40 to 60 aliquot portion of ce or twice repeated The large dose unfits it for delivery in a mixture, Solution, using Pot be dissolved in hot Water by the patient himself —L⁹⁸, 1. 90

The more gel
Lithium Carbonat
U S P include
prepared by dis
Sodium Nitrate
diluting with c
to 5 c c of a 5
produce either
coloration imj
Bromine liber
Water and sh

MAGNESII SULPHAS.

MAGNESIUM SULPHATE.

B.P Syn —EPSOM SALT

$\text{MgSO}_4, 7\text{H}_2\text{O}$, eq. 244 68.

LITHII

aqueous solv FR, SULFATE DE MAGNÉSIUM, GER, MAGNESIUMSULFAT, ITAL, SOLFATO DI MAGNESIO, SPAN, SULFATO MAGNESICO

scaling it

Given

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W

therefore

be kept

in well-closed

bottles or jars

Solubility—1 in 13 of Water, measures 18, 20 in 3 of boiling

Water, insoluble in Alcohol (90 p c)

Medicinal Properties—A mild and safe hydragogue purgative,

operating with little pain or nausea

Used in portal congestion and

chronic constipation and that of lead poisoning, in inflammatory

affections in robust people, in dropsies, and in congestion of brain,

is

go

by reducing blood pressure, it wards off apoplec^{fic} with Ferrous Sulphate it is given in anæmia I = a ingredient in Mistura Alba In the acute form of ei^{tic} on the amœbic variety of dysentery, dr^m doses are gi^{ven}, probably other

When given in conjunction with Diluted Sulphuric may be reduced, the Acid also helps to cover the nause

Successful treatment of tetanus by intraspinal injections of
—L '07, 11 910

Dose—30 to 120 grains = 2 to 8 grammes, for^{es}, given in hot ministration, for a single administration, $\frac{1}{4}$ to $\frac{1}{2}$ oz = 1 grammes

Prescribing Notes—Usually given in solution It has a re^u Swiss and butter taste which is difficult to mask, Sodium Sulphate is much m^{re} It is usually prescribed with Cinnamon Water or Peppermint Water, Chloroform

Mixtures containing Magnesium Sulphate, Phenazone, and a Salic^{id} in down a bulky crystalline deposit, which has been stated to consist of side and Salicylate, but has also been shown to yield a fairly definite per^{cent} of a Magnesium Oxide on ignition—P J '99, 11 332, '02, 1 22, 50, 143

Incompatibles—Potassium and Sodium Carbonates and Bic^{arbonates} Lime Water, Lead Acetate Magnesium Sulphate should not be prescri^{bed} Tartrated Soda, for after some time Magnesium Tartrate will precipit^{ate} following prescription is an example R Sodæ Tartarata, 5j, Magnesⁱⁱ 3ij, Aquæ ad fl 3iss

Official Preparation—Magnesi Sulphas Effervescens Contain^{ing} Mistura Sennæ Composita Used in the preparation of Magnesi C^{omposita} Levis, Magnesi Carbonas Ponderosa, and Liquor Magnesi Carbonatis

Not Official—Eau Saline Purgative, Eau Saline Purgative G^{el} Enema Magnesii Sulphatis, Magma Magnesii, Mistura Salina Laxans, Ma^{gnesii} Benzoas, Magnesi Salicylas and Magnesi Sulphis

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U^S

Tests—Magnesium Sulphate when exposed to warm air lose^s portion of its Water of crystallisation and is converted into a w^{hite} powder At a temperature of 50° to 52° C (122° to 125 6° F) it lo^{ses} one of its seven molecules of Water of crystallisation, equivalent to loss of weight of 7 3 p c At a temperature between 120° to 130° C (248° and 266° F) it loses six molecules of Water, equivalent to a loss of weight of 43 8 p c, and at a temperature of 200° to 230° C (392° to 446° F) it loses the remaining molecule of Water, equivalent to a total loss of weight of 51 1 p c, the salt being rendered anhydrous 244 68 parts of crystallised Magnesium Sulphate yielding 119 52 parts of anhydrous Magnesium Sulphate or 100 parts of the crystalline yield 48 9 parts of the anhydrous salt

It dissolves readily in Water, yielding a solution which is neutral in reaction towards Litmus paper, and which affords the tests distinctive of Magnesium given under that heading Its aqueous solution gives on the addition of Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid It is officially required to indicate 97 36 p c of pure crystallised Magnesium Sulphate, as determined by precipitating the Magnesium as Magnesium Ammonium Phosphate from a solution of 0 5 gramme of the salt in 250 c c of Water,

Dose—60 to 240 grains = 4 to 16 grammes, for administration, for a single administration, $\frac{1}{2}$ to 1 oz = grammes

Foreign Pharmacopœias—U.S., Magnesium Sulphate, Dried Sodium Bicarbonate, 40 g, Dried Tartaric Acid, 21 g crystals, 13.6 g. A granular, effervescent citrate is Official in Belgium and Spain

Not Official

EAU SALINE PURGATIVE (Fr.)—Magnesium Sulphate, 1, Distilled Water, 65. Dissolve and filter

EAU SALINE PURGATIVE GAZEUSE Eau dite de Seville. Dissolve 30 of Magnesium Sulphate and 4 of Sodium Bicarbonate in 6 Fr, (1 oz), filter the solution into a bottle, add 4 of Tartaric Acid in crystals, over. This preparation is also made with 45 and with 60 of Magnesium but when no quantity is indicated, the 30 is above should be used

ENEMA MAGNESII SULPHATIS—Magnesium Sulphate, 15 (5 mm) in Oil, 1, Mucilage of Starch, 15. Dissolve the Magnesium Sulphate in Mucilage, add the Oil and mix.—B.P. 1885, omitted in B.P. 1898. This has been incorporated in the B.P.C.

MAGMA MAGNESIÆ—Magnesium Sulphate, 25, Sodium Hydroxide, 8.1, Water, q.s. to produce 100. Dissolve the Magnesium Sulphate in Water and Sodium Hydroxide in another portion of 400 of Water, and the solutions. Pour the Sodium Hydroxide Solution slowly, in a thin stream into the Magnesium Sulphate Solution, with constant stirring. Allow and precipitate to subside and decant the clear fluid. Wash the Magnesia 3 times with Water by decantation until the washings are free from saline anion. Transfer the Magma to a muslin strainer and allow to drain without pressure. Then re-transfer it to suitable vessels and add sufficient Water to make 100 fluid, and mix thoroughly by stirring. One teaspoonful contains about grains of Magnesium Hydroxide. Average Dose—2 fl drms = 7.1 cc.—U.S.

Note—The Water used in preparing this must be free from organic matter and or the Magma will become discoloured.

A similar preparation with directions closely resembling the above appears in the B.P.C. under the title **Emulsio Magnesiae** (Magnesia Milk), in which Solution of Potassium Hydroxide is used in place of Sodium Hydroxide, and it is about half the strength of Magnesia, there is also a note to the effect that a more concentrated preparation, **Magma Magnesiae** or **Cremor Magnesiae**, may be prepared by doubling the proportion of Magnesium Sulphate which is the strength of the **Magma Magnesiae** of the *U.S.N.F.*, the form given below is incorporated in the B.P.C.

Magnesium Sulphate, 12.50, Solution of Potassium Hydroxide, 114, Distilled Water, q.s. to make 100. General directions are the same as above.

MISTURA SALINA LAXANS—Magnesium Sulphate, 30 grains, Potassium Citrate, 20 grains, Tincture of Hyoscyamus, 15 minims, Chloroform Water, to 1 fl oz.—St. Thomas's.

This has been incorporated in the B.P.C.

MAGNESII BENZOAS $\text{Mg}(\text{C}_6\text{H}_5\text{O})_2 \cdot 3\text{H}_2\text{O}$, eq. 318.08—A white, crystalline powder, soluble 1 in 30 of Water, sparingly in Alcohol (90 p.c.). Introduced as an antipruritic.

Dose—5 to 15 grains = 0.32 to 1 gramme

Tests.—Magnesium Benzoate dissolves in Water, yielding a solution which is neutral or only faintly acid towards Litmus paper. It yields on the addition of Ferric Chloride T.S. a buff coloured precipitate. The concentrated aqueous solution affords when acidified with Hydrochloric Acid a white crystalline precipitate, which, when separated, washed free from mineral acid and carefully dried, should possess the m.p. and answer the tests given under 'Acidum Benzoicum'. The filtrate after the removal of the Benzoic Acid should respond

MANNITOL HEXANITRATE (Hexanitrin) —Odourless, white, needle-like crystals, slightly soluble in Water, soluble in Alcohol and Ether of the hexatomic alcohol, Mannite. It explodes violently on being triturated or struck, and therefore requires great care in handling. Introduced as a vasodilator, stated to possess same action as the Erythrol compound, though not so powerful — *B M J* '95, 11 1213, '98, 1 529, 893

Dose — $1\frac{1}{2}$ to 2 fl drim = 5.4 to 7.1 cc of a 1 p.c. alcoholic Solution

Not Official
MARANTA

ARROW-ROOT

The Starch obtained from the Roots of *Maranta arundinaria*, L., a native of the tropical parts of America and the West Indies, that from Bermuda being considered the best.

A light, white powder, or small pulverulent masses, free from unpleasant odour and taste.

Medicinal Properties Nutrient and demulcent, frequently taken with Milk. It should be first made into a thin paste with cold Milk, and boiling Milk added to make a thick mucilage.

Foreign Pharmacopœias — Official in Mex (Aricoma) and Port (Aiaruta).

Not Official
MASTICHE.

MASTICH

A concrete, resinous exudation, obtained by incisions in the bark of the stem and large branches of *Pistacia Lentiscus*, L., occurring as small, irregular, pale yellow tears, brittle, and either opaque or, far more frequently, transparent. Sp. gr. 1.06 to 1.07.

Produced in the Island of Scio.

Solubility — Insoluble in Water, partly soluble in Alcohol (90 p.c.) and Oil of Turpentine, 2 in 1 of Ether, 2 in 1 of Chloroform.

Medicinal Properties — Used in solution as a temporary stopping for teeth.

Foreign Pharmacopœias — Official in Belg, Norw, Swed (Resina Mastix), Dutch, Hung, Port, Mex and Span (Almaciga) and U.S.

MASTIC DENTAIRE — Mastic 2, Ether 1. Dissolve.

Cotton saturated in this solution is a good stopping for decayed teeth.

MASTIC AND CHLOROFORM — Mastic 2, Chloroform 1. Dissolve. Used for the same purpose as above.

Not Official.
MATICO.

The dried Leaves of *Piper angustifolium*, Ruiz and Pavon. Imported from South America.

Medicinal Properties — An agreeable aromatic astringent, used in all forms of inflammation of the urinary passages, and especially in catarrh of the bladder of the aged. The Volatile Oil has a powerful styptic property, and a solution of it is applied to leech-bites and other small bleeding wounds.

Dose — Of the powder, 30 to 120 grains = 2 to 8 grammes, three times daily.

Foreign Pharmacopœias—Official in Belg, Mex, Port and US Not in the others

Descriptive Notes—The dried leaves as imported from Panama are usually much bent and broken, but in the perfect state are elongate lanceolate, with a rounded, cordate, oblique base, $3\frac{1}{2}$ to 6 inches (9 to 15 cm) long and $1\frac{1}{2}$ to $1\frac{1}{4}$ inches (31 to 37 mm) broad, entire at the margins, with a short leafstalk about $\frac{1}{2}$ inch (12 mm) long, brittle. The veins and veinlets are deeply sunk on the upper surface so as to give a tessellated appearance, and are prominent below, where they form a reticulated network, and densely hairy. The odour is faintly aromatic, and the taste gritty and bitterish. A much larger and broader leaf is occasionally received from Columbia, which possesses a similar surface, but the species that yields it has not been determined. The microscopic features are the short rough hairs, the hypodermal cells of the upper surface, the oil glands and the hypodermal collenchyma above and below the nerve.

The leaves of other species of *Piper*, e.g. (*Piper aduncum*, L., and *P. anisatum*, H. B. and K.) are sometimes mixed with those of true Matico, but neither of these possesses the tessellated nervation of the leaves.

INFUSUM MATICO—Matico Leaves, cut small, 1, boiling Distilled Water, 20. Infuse half an hour and strain.

Dose—1 to 4 fl oz = 28 4 to 113 6 c c

FLUIDEXTRACTUM MATICO—100 of Matico in No. 40 powder is moistened with 80 of a mixture of Alcohol (95 p c) 75 and Water 25, packed in a percolator and macerated for 48 hours, then gradually exhausted with the menstruum. Reserve the first 85, and evaporate the remainder to a soft extract, which dissolve in the reserved portion, and make up with the menstruum to 100.—*U S P*

This has been incorporated in the *B P C*

Official in Mex, 1 in 1

TINCTURA MATICO—Matico Leaves, in coarse powder, 1, Alcohol (60 p c), 5. Macerate 14 days, strain, express, and filter. (1 in 5)

Dose—1 to 2 fl drim = 3 6 to 7 1 c c

Foreign Pharmacopœias—Official in Mex, 1 in 5. Not in the others

Not Official

MEDULLA RUBRA

RED BONE MARROW

The Marrow of ox bones, being a seat of formation of blood corpuscles, has been introduced in the treatment of pernicious anæmia, chlorosis, and hæmoglobinuria. It may be given fresh or raw, spread as a sandwich, also in the form of 'Glycerin Extract,' in gelatin capsules, or as tablets.—*B M J* '94, 1 1172, '95, 1 1084

Red Marrow did not have the slightest effect in three cases of pernicious anæmia, one of which began rapidly to improve on treatment with arsenic.—*L* '96, 1 285

Good results in splenic leucocythemia.—*B M J* '96, 1 840, 956

Fr. has Moelle de Bœuf purifiée, Medulla Bovis depurata

GLYCERIN EXTRACT OF RED BONE-MARROW Veal Marrow, 1, Chloroform Water, 2, Glycerin, 2. Beat up the Marrow with the Glycerin, and add the Chloroform Water, beating the whole together frequently during 1 hour, then strain, and make up to 4 with a mixture of Chloroform Water and Glycerin in equal parts. 4 fluid parts equal 1 part of Marrow.—*P J P*

Extractum Medullæ Rubræ *Syn.* Medullary Glyceride, Glycerin Extract of Red Bone Marrow.—Red Bone Marrow, 25, Chloroform Water, 50, Glycerin, 50.—*B P C*

MEL DEPURATUM.

CLARIFIED HONEY

FR, MIEL BLANC, GER, GERMINGETER HONIG, ITAL, MIEL DEPURATO, SPAN, MIEL DEPURADO

Honey of commerce, melted in a water-bath, and strained, while hot, through flannel previously moistened with warm Water

Medicinal Properties.—Demulcent, laxative, and nutritive, but apt to gripe and occasion flatulence when given in large doses. In the form of **Oxymel** it is a useful addition to gargles and cough-mixtures, as it relieves the pain and dryness of the throat and also dysphagia

Official Preparations.—**Mel Boracis**, **Oxymel**, **Oxymel Scille**. Contained in **Confectio Piperis**

Not Official.—**Aqua Mellis**

Foreign Pharmacopœias.—Official in all, Port, Mellito Simplex, Span, Miel Depurado

Descriptive Notes.—Honey is largely imported from California, Chili and Jamaica, and to a less extent from France, Italy, New Zealand, and other countries. Formerly the white, delicately-flavoured Narbonne Honey imported from France was considered the best. A very white and good-flavoured Honey is now imported from California and Chili. The flavour depends largely upon the flowers from which it is derived, thus heather, linden and clover honey are prepared and sold in Canada. Dark coloured Honey, although often highly flavoured, as is the case with Jamaica Honey, obtains a lower price than paler and weaker-flavoured kinds. During dry summers bees often feed upon the honey-dew or aphid excrement on leaves, which gives it a dark colour, hence, perhaps, the popular prejudice against dark Honey. Pure Honey will always become more or less solid when kept and will contain numerous pollen grains, and by the character of the pollen grains present it is possible to guess at the country from which it was imported. Artificial Honey usually contains glucose prepared from Starch, in the United States 'corn syrup,' prepared from Maize Starch, is used, but such Honey usually contains traces of Sulphuric Acid and of Starch, hence the *P B* test for Sulphates and Starch. Australian Honey, derived largely from species of Eucalypti, has frequently a flavour objectionable to Europeans, Trebizonde Honey, and sometimes North American Honey, has proved to be poisonous or intoxicating, due apparently to the presence of Andromedotoxin derived from flowers of *Rhododendron* and *Lazera*, but these kinds of Honey are rarely met with in English commerce. The term "Virgin" Honey is applied to Honey of light colour produced by a new swarm, or to clear Honey that first runs from the comb of older hives.

Clarified Honey is alone official in the *B P.*, and is directed to be prepared by straining Honey melted in a water-bath through flannel previously moistened with warm Water. In the *U S P* 2 p.c. of paper pulp in shreds is directed to be boiled with the Honey and the

scum which rises removed, the loss incurred being made up with Distilled Water and 5 p c Glycerin by weight added after straining. No sp gr is mentioned, but Vogl states that pure Honey should have a sp gr of 1.410 to 1.445.

The directions for clarification are very necessary for Honey derived from ordinary bee-hives, which may contain debris of immature insects and other impurities, but the Honey derived from frame hives is usually clean and pure, except for pollen derived from bee-bread, which is deposited in some of the cells.

Tests — Clarified Honey is officially required to yield no characteristic reaction for Starch when tested by the Iodine Solution test, the *USP* directs 1 part of Honey to be boiled with 5 of Water, cools the resulting solution and adds Iodine Test solution, when no blue or green coloration should be yielded, the *PG* does not include a test for Starch. It should be free from more than traces of Sulphates as ascertained by dissolving the ash (left on ignition) in Water, acidifying with Nitric Acid and adding Barium Chloride Solution. The *USP* and *PG* test for Sulphates as well as Chlorides in the Clarified Honey diluted with Water. The ash should not exceed 0.25 p c, the *USP* ash limit is 0.3 p c, the *PG* 0.4 p c. The *USP* requires that unclarified Honey, when diluted with twice its weight of Water, should yield a liquid having a sp gr not lower than 1.099, presumably at 25° C (77° F), though in this instance the *USP* does not state so, the *PG* gravity for a similarly prepared liquid is at least 1.111. Clarified Honey is almost invariably levogyrate. It is slightly acid in reaction towards blue Litmus paper. The *PG* requires that it shall contain not more than 0.18 p c w/w of Formic Acid as ascertained by titration with Normal Volumetric Potassium Hydroxide Solution. The *USP* requires that when 1 c c of Absolute Alcohol is carefully run on to the surface of 2 c c of a 25 p c aqueous filtered solution no permanent milky zone should be produced at the junction of the two liquids, indicating the absence of Starch Sugar. The *PG* states that Clarified Honey shall not be rendered turbid by the addition of 2 parts by weight of Alcohol (90 p c), nor should any alteration in colour be manifest when mixed with 1 part by weight of Ammonia Solution. The *USP* includes a test for Cane Sugar with Sulphuric Acid, requiring that when 0.5 c c of a 25 p c aqueous solution is floated carefully on the surface of 2 c c of Sulphuric Acid no coloured line should be immediately produced at the point of contact, the coloration at the end of one hour should amount at most to a yellowish-brown, but no brown or nearly black colour should be developed.

Preparation

OXYMEL OXYMEL

Clarified Honey, liquefied (by weight) 40, Acetic Acid, 5, Distilled Water, *qs* to yield a product of the sp gr 1.320.

Dose — 1 to 2 fl drms = 3.6 to 7.1 c c

Foreign Pharmacopœias — Official in Austr, Clarified Honey, 99, Acetic Acid (96 p c), 1, Dutch, Honey, 19, Acetic Acid (80 p c), 1, Hung, Honey,

50, Acetic Acid (96 p c), 1, Jap, Refined Honey, 8, Acetic Acid, 1, Distilled Water, 1, Port, Honey, 197, Acetic Acid (98 p c), 3, Russ, Honey, 49, Acetic Acid (95 p c), 1, Span, Clarified Honey, 2, White Vinegar, 7, evaporate to 2, Mex, Honey, 100, Acetic Acid, 6 Not in the others

Not Official

AQUA MELLIS—Oil of Bergamot, 2 fl oz, Oil of Lavender, 4 fl drim, Oil of Cloves, 4 fl drim, Oil of Sandal Wood, 1 fl drim, Musk, 10 grains, Tincture of Saffron, 1 fl oz (or q s), Rose Water, 2 pints, Orange Flower Water, 2 pints, Honey, 1 oz, Rectified Spirit, 8 pints.—*Gray's S. 1111*

Oil of Bergamot, 0 75, Oil of Lavender, 0 25, Oil of Cloves 0 25, Oil of Sandal Wood, 0 05, Tincture of Musk, 1 50, Tincture of Saffron, 0 75, Rose Water (undiluted), 16, Orange Flower Water (undiluted), 16 Honey, 0 50, Alcohol q s to produce 100—*B P C*

MENTHÆ PIPERITÆ OLEUM.

OIL OF PEPPERMINT

FR, ESSENCE DE MENTHE POIVRÉE, GR, PEPPERMINT OIL,
ITAL, ESSENZA DI MENTHA, SPAN, ESENCIA DE MENTHA PIPERITA

A volatile oil distilled from fresh flowering Peppermint, *Mentha piperita*, Sm

A clear, colourless, pale yellow or greenish-yellow, oily liquid, possessing a peculiar refreshing odour and a characteristic taste, subsequently producing a sensation of coldness in the mouth The principal constituent of the Oil is Menthol

The Oil official in the *USP* is required to contain not less than 6 p c of Ester calculated as Menthyl Acetate and not less than 50 p c of total Menthol, representing that both free and combined as Ester

It should be kept in well-closed glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light

The variations in quality of the English oils depend (1) upon whether they have been obtained from 'Black Mint' (the ordinary plant) or from 'White Mint', (2) upon the subsequent rectification, so that from the finest double-rectified White Mint to the first crude distillate from the Black Mint there are all manner of gradations, each of them sold as 'Oil Menth Pip Ang'

Dementholised Oil of Peppermint is commonly known as 'Menthene,' and is used for purposes of perfumery. It is pure when it is obtained in a mixture of Ice and Salt, should, on the addition of crystals, set to a more or less solid crystalline mass

American Oil of Peppermint is also the product of *Mentha piperita*, but contains less Menthol

Japanese Oil of Peppermint is obtained from *Mentha arvensis* var *piperita* and contains less Menthol

Solubility—In all proportions of Absolute Alcohol; 2 in 1 (or less) of Alcohol (90 p c), becomes turbid on adding more Alcohol.

Medicinal Properties—An aromatic stomachic and carminative, antiseptic. Allays nausea, relieves spasmodic pains in the stomach. Useful in the flatulent colic of children. Covers the taste of nauseous medicines, such as Rhubarb, and mitigates the

griping effect of purgatives Externally applied it acts as a local anæsthetic and relieves neuralgic pain, *see* also Menthol

Dose — $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Prescribing Notes —*The Oil is taken on sugar, or in pill* See p. 897

Official Preparations —Aqua Menthae Piperitæ and Spiritus Menthae Piperitæ Contained in Pilula Rhei Composita and Tinctura Chloroformi et Morphinæ Composita

Not Official —Essentia Menthae Piperitæ, Syrupus Menthae Piperitæ

Foreign Pharmacopœias —Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital. (Essenza di Menta), Jap. (Oleum Menthae), Mex. (Aceite Volátil de Menta Piperita), Norw. Port. (Essencia de Hortela Pimenta), Russ., Span., Swed., Swiss, and U.S.

The herb is Official in Dan., Fr., Ital., Norw., Swed., and U.S.

Tests —Peppermint Oil has a sp. gr. of 0.900 to 0.920, the official figures are 0.900 to 0.910, the *P.G.* says 0.900 to 0.910, the *USP* 0.894 to 0.914 at 25° C (77° F). It is lavogyrate, the optical rotation being from -20° to -33° in a 100 mm. tube. No figures for optical rotation are given in the *B.P.* It is officially required to dissolve in four times its volume of Alcohol (70 p.c.). The *USP* adds "the solution showing not more than a slight opalescence." The *USP* requires it to also form with an equal volume of Alcohol (94.9 p.c.) a clear solution neutral in reaction towards Litmus paper. The *B.P.* requires that when a portion of the Oil is cooled to -8.3° C (17° F) and a few crystals of Menthol are added, a considerable separation of Menthol should take place. This test has been adversely criticised and its deletion from the Pharmacopœia has been recommended. The *B.P.* does not require a definite yield of Menthol nor suggest a method of determination. It has been recommended that the Menthol, both free and combined, should be determined by the usual methods.

The *USP* requires that the Oil shall contain not less than 6 p.c. of Ester calculated as Menthyl Acetate, and not less than 50 p.c. of total Menthol (both free and combined as Ester), as determined by weighing 10 c.c. of the oil in a flask and saponifying it by boiling under a reflux condenser with 25 c.c. of Semi-normal Volumetric Potassium Hydroxide Solution, and titrating the excess of Volumetric Hydroxide Solution, with Semi-normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. The number of c.c. of Semi-normal Sulphuric Acid Solution required, subtracted from the 25 c.c. of Semi-normal Potassium Hydroxide Solution used and the difference first multiplied by 9.834 and then divided by the weight of the Oil yields the percentage w/w of Menthyl Acetate present in the sample. The Oil remaining after saponification is then washed repeatedly with Water and transferred to an acetylation flask, and the Menthol converted into an Acetyl derivative by boiling for one hour with 10 c.c. of Acetic Acid Anhydride and about 1 gramme of anhydrous Sodium Acetate. It is allowed to cool, freed from excess of Acid by washing first with Water and then with Sodium Hydroxide T.S. until the mixture is slightly alkaline.

in reaction towards Phenolphthalein Solution and dried by fused Calcium Chloride

A measured quantity of 5 c.c. of the acetylated product is transferred to a flask and its weight accurately recorded. It is then saponified for one hour under a reflux condenser, with 50 c.c. of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution, the excess of Volumetric Alkali being titrated with Semi-normal Volumetric Sulphuric Acid Solution, Phenolphthalein Solution being employed as an indicator of neutrality. The number of c.c. of Volumetric Acid Solution required is subtracted from 50, the difference is multiplied by 7.749 and the product divided by the weight of acetylated product employed for the determination less the difference (between the number of c.c. of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution employed for the determination and the number of c.c. of Semi-normal Volumetric Sulphuric Acid Solution required to neutralise the excess of Volumetric Alkali Solution multiplied by 0.021), the quotient will represent the % of Menthol present in the Oil of Peppermint. The percentage of combined Menthol varies from 3 to 14 p.c. and the total Menthol from 30 to 70 p.c. In addition to a determination of the combined and total Menthol, a determination of the amount of Menthone is often useful. It may be carried out on a separate portion of the saponified Oil by diluting it with twice its volume of Alcohol and boiling it with metallic Sodium. The Menthone is reduced to Menthol, and the amount of Menthol produced may be determined by acetylation.

The Oil of *Mentha piperita* is, as a rule, distinguished from that of *Mentha pulegiensis* by developing a blue colour or red fluorescence when mixed with 4 volumes of Glacial Acetic Acid, this colour is not developed if air be excluded, and, depending as it does upon some minor constituent destroyed by prolonged exposure to sunlight, it may not be given by some old samples. Two varieties of the Oil, black and white Peppermint respectively, are grown in and about Mitcham. It is stated (*P. J.* '96, 1 125) that the white Oil of Peppermint may be distinguished from the black by its having greater optical activity, and not depositing Menthol at a low temperature, and containing a greater proportion of Esters of Menthol, and in giving an intense blue coloration, with red fluorescence, with Glacial Acetic Acid.

The more generally occurring sophistications of the Oil are abstraction of Menthol, adulteration with Turpentine Oil, Cedar Wood Oil, and the addition of other compound esters. The presence of dementholated Oil is indicated by the decrease in the sp. gr. and optical rotation and by the diminution in the total Menthol content. Turpentine Oil may be detected by its effect on the optical rotation, and by the optical rotation of the various fractions obtained on distilling the Oil. It may also be detected by its effect on the solubility of the Oil in Alcohol (70 p.c.). The figures given by Parry and Bennett in an examination of some Peppermint Oils (*C. D.* '04, 1 854) clearly point to adulteration of the Oil with a substance having the nature of a sesquiterpene, which led them to the opinion that the adulterant was Cedar Wood Oil. The Oils were insoluble in

Alcohol (70 p c), and the later fractions of the various distillates, in addition to being much less soluble than the distillate from pure Peppermint Oil, had in some cases a distinct taste of Cedar Wood Oil.

The presence of Acetin, a Glycerin Acetic ester, has been detected as an adulterant by Bennett (*C D* '03, i 591). Oils containing the adulterant are distinguished by a higher sp gr, the decrease in the optical rotation, the insolubility in Alcohol (70 p c), and the exceptional high ester figure obtained in the determination of the combined Menthol. A comparison of the distillates obtained on fractionating the Oil also readily reveals the presence of the adulterant. The production of the odour of Ethyl Acetate in the cold on the addition of Potassium Hydroxide is suggested as a characteristic, but hardly conclusive, test. Parry and Bennett (*C D* '03, ii 154) have detected the presence of African Copaiba Oil in some adulterated samples of Peppermint Oil, by the isolation of a fraction of high boiling point, which contained a body having a sp gr within the limits of Peppermint Oil, but with a strongly positive optical rotation and a high refractive index, which, from its physical characters and chemical reactions, was a substance belonging to the sesquiterpene series. Messrs Schimmel and Co have not met with Oils containing this adulteration. The Oil consists principally of Menthol, together with Acetic and Valerianic Acid esters of Menthol, Menthone, 2 lœvogyrate terpenes, a dextrogyrate sesquiterpene and Phellandrene.

Alcohol—It should be soluble in 4 volumes of Alcohol (70 p c), *BP* and *USP*, in 4 to 5 parts by weight of Alcohol (68 to 69 p c), *P U*. The solution should not show more than a slight opalescence, *USP*.

Optical Rotation—It is lœvogyrate, the angle of rotation varying from -25° to -33° in a 100 mm tube at 25°C (77°F), *USP*.

Mercuric Chloride—If from 25 c c of the Oil the 1st 1 c c of distillate be collected and poured on an aqueous Mercuric Chloride Solution, a white film should not form at the zone of contact after a short time (absence of Dimethyl sulphide found in non rectified oils), *USP*.

Preparations

AQUA MENTHÆ PIPERITÆ PEPPERMINT WATER

Oil of Peppermint, 77 minims, Water, $1\frac{1}{2}$ galls, distil $\frac{1}{2}$;
(Oil about 1 in 1000)

Dose—1 to 2 fl oz = 28.4 to 56.8 c c

Foreign Pharmacopœias—Official in Belg, Spirit of Peppermint, 30 in 1000, Dan and Russ, 1 of Oil in 2000, *US*, 1 of Oil in 500, Austr and Hung, 1 of fresh leaves to produce 5 of distillate, Dutch, Ger, Swed and Swiss, 1 to produce 10, Fl, Poit and Span, 1 to produce 1, Ital, 1 to produce 2, Jap, 1 to produce 30, Mex, 1 of fresh plant to produce 4.

SPIRITUS MENTHÆ PIPERITÆ SPIRIT OF PEPPERMINT

Oil of Peppermint, 1, Alcohol (90 p c), *qs* to yield 10

BP '85 was 1 in 50

Dose—5 to 20 minims = 0.3 to 1.2 c c

Foreign Pharmacopœias—Official in Austr, Oil, 1, Alcohol (90 p c), 19, Belg (*Spiritus Menthæ*), Oil, 1, Alcohol, 99, Fr (*Teinture d'Essence de Menthe*), Oil, 2, Alcohol, 98, Ger and Jap (*Spiritus*

Menthæ, 1 in 10, Swiss, 3 Oil in 100, U S, from the leaves and oil, about 1 in 10, Mex (Alcoholatura de Menta), 1 of fresh plant macerated in 3 of Alcohol (80 p c) for 8 days, Span (Alcohol de Menta Piperitæ), 10 of the fresh leaves and tops macerated in 20 of Alcohol (60 p c) for 2 days, distil 10 Not in the others

Not Official

ESSENTIA MENTHÆ PIPERITÆ —Oil of Peppermint, 1, Rectified Spirit, 4 —*B P* 1885, omitted in *B P* 1898, but now incorporated in the *B P C*

SYRUPUS MENTHÆ PIPERITÆ —Spirit of Peppermint, 1, Simple Syrup, *q s* to yield 8

A good flavouring for nauseous medicines

Foreign Pharmacopœias —Official in Austri, Water, 2, Sug, Spirit of Peppermint, 3, Syrup, 97, Ger Peppermint Leaves, 2, Alcohol, 1, Distill Water, 10, macerate for 24 hours, press, filter, and in 7 of the filtrate dissolve 13 of Sugar to make 20 of Syrup by weight

MENTHÆ VIRIDIS OLEUM.**OIL OF SPEARMINT**

NO Syn —**MENTHÆ CRISPÆ OLEUM**

FR, HUILE VOLATILE DE MENTHE VERBE, GER, ROMISCHMINTÖL, ITAL, ESSENZA DI MENTA, SPAN, ESNCIA DE MENTA

A colourless, pale-yellow, or greenish-yellow, limpid liquid, having a characteristic odour and taste, distilled from fresh flowering Spearmint, *Mentha viridis*, L

The principal constituent of the oil is Carvone, it also contains Linalool, lavo-limonene and possibly lavo-pinene. It is liable to become darker in colour with age and exposure to light and air, and should therefore be kept in well-stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as much as possible from the light and from contact with the air

Solubility —In all proportions of Absolute Alcohol, 1 in 1 (*or less*) of Alcohol (90 p c), becomes milky on adding more Alcohol

Medicinal Properties. —Similar to those of Oleum Menthæ Piperitæ

Dose — $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c

Prescribing Notes —The oil is given on sugar, or made into pills with Liquorice Powder and Soap See p 897

Official Preparation —Aqua Menthæ Viridis.

Not Official. —Spiritus Menthæ Viridis

Foreign Pharmacopœias —Official in Hung, Port (Essencia de Hortela), Russ and U S (sp gr 0.914 to 0.934 at 25° C (77° F)). Not in the others

Tests —It is officially required to have a sp gr of 0.920 to 0.940; the *USP* says 0.914 to 0.934 at 25° C (77° F). It is levogyrate, possessing an optical rotation from 35° to 48° in a tube of 100 mm, it should form a clear solution when mixed with an equal volume of a mixture of equal parts of Absolute Alcohol and Alcohol (90 p c) The *USP* states that it should form a clear solution when mixed

with an equal volume of Alcohol (80 p c), the solution becoming turbid upon further dilution. The Oil is not official in the German Pharmacopœia

AQUA MENTHÆ VIRIDIS SPEARMINT WATER

Oil of Spearmint, 77 minims, Water, 1½ galls, distil ¾
(Oil about 1 in 1000)

Dose —1 to 2 fl oz = 28 4 to 56 8 c c

Official in U S, 1 in 500, Port (Aguado Hortela)

Not Official

SPIRITUS MENTHÆ VIRIDIS —Spearmint, 1, Oil of Spearmint, 10, Alcohol (95 p c), to yield 100, macerate for 24 hours and filter —*U S P*

Oil of Spearmint, 1, Alcohol (90 p c), to make 10 —*B P C*

MENTHOL.

MENTHOL

C₁₀H₂₀O, eq 154 98

FR, **MENTHOL**, **GFR**, **MLNTHOL**, **ITAL**, **MENTHOL**, **SPAN**, **MENTOL**

Large, colourless, acicular, or prismatic crystals, obtained from the Volatile Oil distilled from the fresh Herb of *Mentha arvensis*, L, var *piperascens et glabrata*, Holmes, and of *Mentha piperita*, Sm

It possesses the strong characteristic odour and taste of Peppermint, subsequently producing a sensation of warmth on the tongue, and upon inspiration of air a sensation of coldness

It should be kept in well stoppered glass bottles in a cool atmosphere

Solubility —Almost insoluble in Water and Glycerin, soluble 5 in 1 of Alcohol (90 p c), 4 in 1 (nearly) of Chloroform, 8 in 3 of Ether, 10 in 7 of Petroleum Spirit, 1 in 4 of Olive Oil

Menthol forms a liquid when rubbed with equal parts of either Carbolic Acid, Chloral Hydrate, or Thymol, 3 of Menthol and 2 Camphor form a liquid at ordinary temperatures, but when in equal parts is liquid only whilst warmed

Medicinal Properties —Antiseptic, stimulant, carminative, local anæsthetic. Applied externally as a local analgesic in some forms of neuralgia and headache, also in rheumatism, in pruritus and in pleurodynia and toothache

Used as a **snuff**, along with Boric Acid 2 parts, and Ammonium Chloride 3 parts, also dissolved in oil as a spray for influenza, hay-fever, coryza and ozæna

Menthol and Eucalyptus Oil dissolved in Alcohol (90 p c) is used with an oro nasal inhaler for cold in the head,

10 minims of a 20 p c alcohol solution of Menthol used for an hour or so on the sponge of an inhaler is useful in the troublesome cough of phthisis —*Edin Med Jour* '05, 465

A 20 p c solution in Olive Oil (with 3 p c Guaiacol) as an intralaryngeal injection (20 to 30 minims) in phthisis and bronchiectasis —*Pr* lvi 276

Intralaryngeal injection of a 10 p c solution in Olive Oil, along with Boric Acid and Quinine Sulphate internally in gangrene of the lung —*B M J* '99, i 71

A good remedy in painful enteritis with mucous diarrhoea — *M J B* '05, 239, and in gastralgia and vomiting — *B M J B* '07, 1 36, *P* '07, 1 717

Spray containing 5 to 20 pc of Menthol recommended in tubercular laryngitis — *T G* '87, 762

Menthol and Iodoform equal parts as a surgical dressing — *B M J* '88, 1 933

Dose — $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Prescribing Notes — *It is best made into pills by the addition of Soap and Dispensing Syrup. Usually used in the form of cones and pencils, also by insufflation, or as an ointment, pigment or plaster*

Pastilles may be obtained containing Menthol $\frac{1}{10}$ or $\frac{1}{6}$ grain in each, also Menthol and Cocaine $\frac{1}{8}$ grain of each, Menthol $\frac{1}{10}$, Extract of Rhatany, 2 grains, and Bromide of Ammonium, 1 grain, Menthol, $\frac{3}{10}$ grain, and Menthol, $\frac{1}{10}$ grain, and Caffein, $\frac{1}{4}$ grain, Menthol, $\frac{3}{10}$, Cocaine, $\frac{1}{10}$ grain, and Red Gum, 2 grains, Menthol, $\frac{1}{10}$ grain, Eucalyptus, 1 minum, and Cocaine, $\frac{1}{10}$ grain, Menthol, $\frac{1}{10}$, and Heroin, $\frac{1}{20}$ grain. Menthol, $\frac{1}{10}$, and Terpine, $\frac{1}{4}$ grain. Tablets containing $\frac{1}{16}$, $\frac{1}{20}$, $\frac{1}{4}$, $\frac{1}{2}$ grain, also in combination with other drugs

Official Preparation — *Emplastrum Menthol*

Not Official — *Applicatio Menthol, Aqua Menthol, Gossypium Menthol, Insufflatio Menthol, Insufflatio Menthol Composita, Nebula Menthol, Nebula Menthol Composita, Nebula Menthol cum Cocaina, Pastillus Menthol, Pigmentum Menthol, Pigmentum Menthol, Mentholacea Vasolumentum Mentholi, Parogenum Menthol, Coryza, Unguentum Menthol, Menthol Valerianate, Menthol*

Foreign Pharmacopœias — Official in Austr, Belg, Dan, Dutch, Fr, Ger, Jap, Norw, Russ, Swed, Span and Swiss (Mentholum), Ital (Mentolo), Mex (Mentol), US (Menthol) Not in the others

Tests. — Menthol when pure has a mp from 43.5° to 45° C (110.3° to 112.1° F), the *BP* gives the mp as 42° C (107.6° F) and that it should not exceed 43° C (109.4° F)

Tvisi has shown that commercial Menthol melts at 39.02° C (102.23° F), dried Menthol at 41.57° C (106.82° F) and purified Menthol 42.78° C (109° F), and is of opinion (*YBP* '00, 453) that only dried and purified Menthol agrees with the Pharmacopœia requirements, the *USP* and the *PG* both give 43° C (109.4° F). It boils at about 217° C (422.6° F) when the mercury thread of the thermometer is immersed entirely in the vapour, the *BP* does not refer to the boiling point, both *USP* and *PG* give 212° C (413.6° F). It dissolves readily in Alcohol, forming a clear solution which should be neutral in reaction towards Litmus paper, and which is lavogyrate. It volatilises at ordinary temperatures and rapidly on heating, the Pharmacopœia states that when boiled with Sulphuric Acid diluted with half its volume of Water it acquires a brownish blue or ultramarine colour, the acid becoming brown, this colour reaction is considered (*YBP* '00, 338) of extremely doubtful value.

The *PG* states that it yields with 10 parts (by weight) of Sulphuric Acid a brownish-red, turbid fluid which clears in the course of a day, and shows on its surface a colourless layer which no longer possesses an odour of Menthol. No method of determination is given in either of the three Pharmacopœias, but it may be determined

as described under *Oleum Menthæ Pipentæ* by acetylation with Acetic Anhydride

The more generally occurring impurities are Wax, Paraffin, or inorganic substances. All three substances may be immediately detected by any residue remaining after the volatilisation of the Menthol. The *USP* and the *PG* include a test for Thymol, requiring that no green coloration shall be produced when a few drops of Sulphuric Acid containing a drop of Nitric Acid are added to a solution of a few crystals of Menthol in Glacial Acetic Acid. The Pharmacopœia states that Menthol crystals are usually more or less moist from adhering oil, and Schimmel and Co have pointed out in their *Semi-Annual Report* for October 1907 that by this statement the *BP* admits an adulterated article.

Heat—It volatilises slowly at ordinary temperatures, *USP*. It is completely volatilised when heated at the temperature of a water bath, *BP*. 0.1 gramme should not leave a weighable residue when so heated, *PG*. Heated in an open dish it should volatilise without residue, *USP*.

Sulphuric Acid—If Menthol be boiled with a mixture of Sulphuric Acid 2 volumes and Water 1 volume, the Menthol becomes indigo blue or ultramarine in colour, and the Acid turns brown, *BP*. It gives with 40 parts of Sulphuric Acid a brownish red turbid liquid which clarifies in the course of a day, and on its surface shows a layer which does not smell of Menthol, *PG*.

Sulphur, Nitric and Acetic Acids—If Menthol be added to a mixture of Acetic Acid 1 c.c., Sulphuric Acid 6 drops, and Nitric Acid 1 drop, no coloration should appear, *PG*. 3 drops of Sulphuric Acid and 1 drop of Nitric Acid added to a solution of a few crystals of Menthol in 1 c.c. of Glacial Acetic Acid should produce no green colour, *USP*.

Preparation

EMPLASTRUM MENTHOL MENTHOL PLASTER

Menthol, 1½, Yellow Beeswax, 1, Resin, 7½. Mix in the Menthol when the melted Resin and Beeswax have cooled to 160° or 170° F (71.1° or 76.7° C).

The quantity of Menthol is reduced and that of the Resin slightly increased to that of *BP* 1885.

Not Official

APPLICATIO MENTHOL—Menthol, 2, Chloroform, 8, Pure Ether (sp gr 0.720), 8, Eau de Cologne, 4. A good external application for neuralgia.

AQUA MENTHOL—Menthol, 8 grains, Alcohol (90 p.c.), 2 fl. dr. , Distilled Water, 20 fl. oz.—*Bournemouth Formulary*
Menthol, 0.10, Alcohol, 0.15, Distilled Water, 100.—*BP C*

GOSSYPIMUM MENTHOL—Menthol, 7 grains, White Adipose Oil, 3 minims, Pure Ether, 6 fl. dr. , Cotton Wool, in a thin sheet, 60 grains.

INSUFFLATIO MENTHOL (Nasal)—Menthol, in powder, 5 grains, Bismuth Oxychloride, ½ oz., Dried Starch, in fine powder, to 1 oz.

INSUFFLATIO MENTHOL COMPOSITA—Menthol, 2 dr. , Ammonium Chloride, 3 dr. , Boric Acid, to 1 oz.—*Central Throat*

Pulvis Menthol Compositus—Menthol, 2, Boric Acid, in powder, 3, Ammonium Chloride, in powder, 8.—*St Thomas's*

For insufflation

Insufflatio Mentholis—Menthol, in powder, 5, Ammonium Chloride, in fine powder, 45, Boric Acid, in powder, 50.—*BP C*

Insufflatio Menthol *Syn* Menthol Snuff—Menthol, 1, Ammonium Chloride, 8, Boric Acid, 2, Lycopodium, 6.—*Martindale*

Almost as ill adapted for a hypnotic as Ether—*L* '99, ii 5

Dose.—15 to 60 minims (in Water) = 0.9 to 3.6 cc

Not Official

METHYLENE BLUE

$C_{16}H_{18}N_3SCl$, eq 317.39

TETRAMETHYLTHIONINE HYDROCHLORIDE, METHYLTHIONINE HYDROCHLORIDUM,
METHYLTHIONINE HYDROCHLORIDE

Prismatic crystals having a bronze-green fluorescence or as a dark green crystalline powder. It stains the skin an intense blue. It dissolves readily in Water, forming an intensely blue coloured solution.

It may be converted into Hydrogen Sulphide on an oxidation product of . . .

Medicinal Properties.—It has been employed as an analgesic in rheumatism and sciatica, also in migraine, in some cases it may produce gastric irritation and cystitis—*B M J* '98, ii 1055, *B M J E* '00, ii 75. It acts specifically in malaria and also in gonorrhoea—*B M J E* '01, i 104, *T G* '00, 474.

Intramuscular injection for diagnosing the degree of penetrability of the kidney tissue. 1 cc of a 5 p.c. solution injected deeply into the gluteal muscles. Urine should be greenish in colour half an hour after injection. A more rapid appearance of the colour, or its shortened elimination and delayed excretion, serve as criteria for the degree of the renal lesion—*T G* '00, 404, *Merck's Archives*, '99, 104.

Use in malaria further advocated (*Pr* lxxviii 682), 2-grain doses with powdered nutmeg and a small quantity of Codeine to prevent strangury and nausea.

Dose.—1 to 5 grains = 0.06 to 0.32 gramme

Prescribing Notes.—It is conveniently given in pills, using 'Diluted Glucose' as the excipient or in capsules. Care must be taken to ensure purity of the sample, and only Zinc-free Methylene Blue should be used for internal administration. Methylene Blue is frequently prescribed in capsules with Oil of Sandal Wood.

Foreign Pharmacopœias.—Official in Fr (Bleu de Méthylène officinale), Tex (Azul de Metilena), Swiss (Methylenum coeruleum), U.S. (Methylthionine Hydrochloridum).

Tests.—Methylene Blue yields, when dissolved in Water, a very deep blue coloured solution, changing on the addition of Hydrochloric Acid to a lighter shade of blue, and on the addition of Sodium Hydroxide Solution to a purplish shade. When the latter reagent is added in excess a dirty violet precipitate is produced. When treated with Zinc and . . . is converted . . . The double Zinc Salt (Tetramethylthionine . . . is also known commercially under the name of Methylene Blue, and it is necessary, therefore, to distinguish between this and Methylene Blue for medicinal use. Medicinal Methylene Blue should not leave more than 0.4 p.c. of ash which, when dissolved in Hydrochloric Acid should not give a . . . destructive of Zinc given under that heading. It should not afford any reaction for Arsenic when ignited with dry Sodium Carbonate and Potassium Nitrate and examined by the modified Gutzeit's test.

Antirheumatin, a mixture of Sodium Salicylate and Methylene Blue, introduced as an antirheumatic, and administered internally in doses of 1 to 2 grains = 0.06 to 0.12 gramme. It must not be confounded with Antirheumin, which is a preparation containing Fluorine compounds.

METHYL VIOLET --Under the fancy name **Pyocetanin** (blue) it has been recommended in the internal and local treatment of malignant tumours. *T G* '94, 706, *B M J E* '94, 11 12 Locally in corneal ulceration --*T G* '93, 55

MEZEREI CORTEX

MEZEREON BARK

FR, MI ZERLON OU BOIS CHNTIL, GER, SHIDIFASERINDI, LEAF, MI ZERRO,
SIAN, MI CERION

The dried Bark of *Daphne Mezereum*, L, of *Daphne Laureola*, L,
or of *Daphne Gnidium*, L

Medicinal Properties --Externally, rubefacient and vesicant
Rarely given alone internally as a gastric stimulant, but it appears as
an ingredient in *Liquor Sarsæ Compositus Concentratus*

Official Preparation --Used in the preparation of *Liquor Sarsæ Compositus Concentratus*

Not Official --*Extractum Mezerei Æthereum* and *Unguentum Mezerei*

Foreign Pharmacopœias --Official in Jap, Mex (*Mezereon*), Port
(*Tlovisco*), Swed, Swiss and U S Not in the others

Descriptive Notes --Mezereon Bark as met with in commerce consists of loose, more or less quilled, strips of thin bark about $\frac{1}{2}$ in thick, finely fibrous, and too tough to be broken, the colour varies from pale, dull greenish-brown in that of *D Laureola* to reddish-brown in that of *D Mezereum* and dark purplish in that of *D Gnidium*. Mezereon Bark is chiefly imported from Germany, but the last-named from Algeria and the South of France. In *D Laureola* the leaf scars are crowded at intervals, but in the two latter are irregularly scattered, and the bark of *D Gnidium* is hairy towards the apex. The outer corky coat readily separates from the green fibrous layer, which is white and satiny on the inner surface. The bark has an acid taste and but little odour. It apparently owes its activity to a resin. The official bark is referred to the three species above named, but to *D Mezereum* and other European species in the *U S P* and the distinguishing feature is there mentioned that the transverse section exhibits numerous groups of bast fibres in the secondary bast. Mezereon Bark is not used in powder, but the principal microscopical features are the very long fibres, often 3 mm long and only 5 to 10 μ in diameter, and the thin walled tubular cells of the cork hexagonal in tangential section.

Tests --Mezereon Bark contains from 3 to 4 per cent of ash, which latter figure should not be exceeded.

Not Official

EXTRACTUM MEZEREI ÆTHEREUM (*B P* '95) --The Æther soluble portion of alcoholic extract of Mezereon Bark.

This has been incorporated in the *B P C*

Foreign Pharmacopœias Official in Port (*Extracto de Tlovisco*)

with Alcohol only, U S, Fluid Extract, 1 in 1, Mezereon Bark treated with a mixture of Alcohol (90 p c), 4, Water, 1 Not in the others

UNGUENTUM MEZEREI—Ext Garou, 4, Lard, 90, White Wax, 10, Alcohol, 9

Not Official MORI SUCCUS

MULBERRY JUICE

The deep purple juice of the ripe Fruit of *Morus nigra*, L. Sp gr about 1 060

Medicinal Properties—Refrigerant and laxative, serves to prepare a grateful drink in febrile cases, and as a flavouring and colouring agent

Foreign Pharmacopœias—Official in Fr, Suc de Mûre, Mex, Jugo de Moras, Port, Amorias, Span, Zumo de Moras

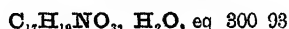
The Fruit is official in Ital

SYRUPUS MORI—Mulberry Juice, 20, Refined Sugar, 36, Alcohol (90 p c), 2½, heat the Juice to the boilingpoint, and, when it has cooled, filter it, dissolve the Sugar in the filtered liquid by a gentle heat, and add the Alcohol, the product should weigh 54 Sp gr 1 330

Dose—1 fl drm = 3 6 c c

Foreign Pharmacopœias—Official in Austr, Belg, Fr (Sirope de Mûres), Hung, Ital and Swiss Not in the others

Not Official MORPHINA



Colourless, shining, rhombic prisms, or as a white, odourless, crystalline powder, having a bitter taste and an alkaline reaction It is the principal alkaloid obtained from Opium

Solubility—1 in 1000 of cold Water, 1 in 100 of Alcohol (90 p c), 1 in 10 of Oleic Acid, 1 in 125 of Glycerin, but the solubilities depend very largely on the physical condition of the alkaloid Insoluble in Ether (thus differing from Narcotine) Aqueous alkalis, even Lime Water, dissolve it readily when freshly precipitated, Animonium however, but sparingly, where a very strong solution is required Hypophosphorous Acid has been suggested as a solvent

Medicinal Properties (see Morphine Hydrochloridum)—Owing to its slight solubility in Water it is rarely given in its purely alkaloidal form

Dose— $\frac{1}{16}$ to $\frac{1}{2}$ grain = 0 0067 to 0 032 gramme

Official Preparations—Morphine Acetate, Morphine Hydrochloride, and Morphine Tartrate

Not Official—Elixir Acetomorphine et Terebinthinae, Tinctur Heroin cum Terpene, Glycerinum Acetomorphine, Glycerinum Heroin, Compositum, Linctus Acetomorphine, Linctus Heroin, Pastilla Morphi, Pastilla Morphi Composita, Pastilla Morphi, Diacetyl-morphine (Heroin), Diacetyl-morphine Hydrochloride, Benzyl-morphine Hydrochloride (Peronine), Morphinum Hydrochloride (Dionine), Morphine Hydrobromide, Morphine Lactate, and Morphine Sulphate, Liquor Morphine Sulphatis, Pulvis Morphine Compositus

Foreign Pharmacopœias—Official in Fr, Hung, Mex (Morfina), Port, Span and U S Not in the others

Tests.—Morphine when heated to 100° C (212° F) loses its Water of crystallisation slowly, but when dried at 110° C (230° F) the Water of crystallisation is rapidly lost It melts at about 230° C (446° F), and at a somewhat higher temperature the alkaloid turns brown It is only very slightly soluble in

Water and the aqueous solution has an alkaline reaction towards red Litmus paper.

A particle of solid Morphine yields the following characteristic colour reactions when treated with a drop of a perfectly neutral Ferric Chloride T S, or with Ferric Ammonium Sulphate Solution (10 p.c. w/v), it yields a very characteristic deep greenish blue colour, changing to green on adding an excess of the reagent, with Sulphuric Acid it yields no coloration, or at the most but a slight yellowish tint, but if heated on the water bath it assumes a brownish coloration. Dott has pointed out that solid Morphine yields a distinct though faint pink colour when treated with Sulphuric Acid, if heated with Sulphuric Acid to 150°C (302°F) a dirty green or rose red colour is produced and if the temperature be still further raised the solution becomes almost black, when cooled and diluted with Water the solution yields a greenish blue colour, which on the addition of Ammonia Solution changes to green, when heated on a water-bath for 10 or 15 minutes with Sulphuric Acid, cooled and a few drops of diluted Nitric Acid added, a violet coloration, changing rapidly to blood red, is produced, Sulphuric Acid, to which a small quantity of Sodium Arsenate has been added affords a bluish green coloration, which on raising the temperature passes from green to deep blue and finally to a dark olive green, if a small quantity of Bismuth Oxynitrate be added to a mixture of Sulphuric Acid and the alkaloid, a purplish brown coloration is yielded. Sulphuric Acid containing 0.5 p.c. of Ammonium Molybdate (Froehde's) reagent yields with Morphine a violet blue coloration, changing to green and ultimately to deep blue. Sulphuric Acid containing a crystal of Potassium Iodate yields a dark brown coloration, Sulphuric Acid containing in each c.c. a drop of Formaldehyde Solution yields an intense purple coloration, a similar purple coloration is produced if the alkaloid be mixed with 2 parts of Cane Sugar, and if a drop of concentrated Sulphuric Acid be added, the colour changing gradually from blood red to brownish red and becoming brown on dilution with Water. Sulphuric Acid containing a crystal of Potassium Bichromate is slowly reduced with the production of a green coloration. Nitric Acid yields an orange red colour changing to yellow. With Potassium Ferricyanide Solution containing a drop of neutral Ferric Chloride T S it yields a deep blue coloration. It reduces Iodic Acid with liberation of Iodine, and if solution of Starch be present a blue coloration of Iodide of Starch is immediately produced. In employing this test it is essential that the reagent itself should not give free Iodine on treatment with a drop of diluted Sulphuric or Acetic Acid. It dissolves readily in diluted Hydrochloric or Sulphuric Acid, and if the alkaloid be carefully and exactly neutralised and the solution be of a sufficient degree of concentration, it will respond to the following tests. Ammonia Solution produces a white precipitate soluble with difficulty in excess of the reagent, Ammonium Carbonate Solution and Solution of Lime Water also produce a white precipitate speedily becoming crystalline. The alkali Carbonates in excess have a tendency to redissolve the precipitate, but it is insoluble in excess of the Bicarbonates. Potassium or Sodium Hydroxide Solution yields a white precipitate readily soluble in excess. Mercuric Potassium Iodide (Mayer's) Solution produces a white gelatinous precipitate. Potassium Ferricyanide Solution containing 1 or 2 drops of Ferric Chloride T S produces in solutions faintly acidified with Hydrochloric Acid a blue coloration or a precipitate of Prussian blue. Morphine is alkaline in reaction towards red Litmus, Methyl Orange, and Iodeosin Solutions, it forms salts which are neutral in reaction towards these indicators. It may therefore be titrated with Normal or Tenth normal Volumetric Hydrochloric or Sulphuric Acid Solution, using one or other of these solutions as an indicator of neutrality. Methyl Orange or Iodeosin Solution is most suitable for the purpose, a known weight of the alkaloid may be dissolved in an excess of Normal Volumetric Sulphuric Acid Solution, a few drops of Methyl Orange or Iodeosin Solution added and the excess of Acid titrated with Normal Volumetric Sodium Hydroxide Solution. The number of c.c. of volumetric alkali solution in excess is subtracted from the number of c.c. of volumetric acid solution used, the difference representing the number of c.c. of normal volumetric acid solution neutralised by the alkaloid. 1 c.c. of Normal Volumetric Sulphuric Acid Solution represents 0.30093 grammes of hydrated Morphine or 0.28305 gramme of anhydrous Morphine. Morphine may be distinguished from

Codeine by the Nitric Acid test, Codeine yielding a red, by Ferric Chloride T.S., which produces a dull greenish blue coloration, but which does not affect Codeine, by Sulphuric Acid, containing a trace of Selenium Acid, which gives with Morphine a blue coloration changing to green and finally to brown, and with Codeine a green coloration changing to blue and afterwards to a grass green, this test also serves to distinguish Morphine from Narcotine, the latter giving a green coloration changing to brown to cherry-red colour test, Bichromate test, the former yielding with these reagents a purple coloration, the latter a deep crimson colour.

The more generally occurring impurities are other alkaloids, *e.g.*, Narceine, Narcotine, Thebaine, and Pseudomorphine, Ammonium salts, Meconic Acid or Meconates, and mineral matter.

The Morphine obtained from any of the official Morphine salts by precipitation is required by the *BP* to yield little or nothing to Benzol, which is considered sufficient evidence of the absence of other alkaloids, the *USP* requires that 0.2 of a gramme of Morphine should yield a clear solution with 4 c.c. of Potassium Hydroxide T.S., and that no insoluble residue should remain, the Solution of Ammonia indicating the absence of Ammonium salts. If a few drops of Ferric Chloride T.S. be added to a solution of 0.1 of a gramme of Morphine in 10 c.c. of diluted Hydrochloric Acid no red coloration should be produced, indicating the absence of Meconic Acid or Meconates. When ignited with free access of air, Morphine should leave no weighable residue, indicating the absence of mineral matter.

A reference to a process of separating small quantities of Morphine from Tincture and such like preparations is mentioned in the large type under Tincture Compositus, a reference is also made under the same heading to the association, in cough mixtures, of Morphine with the alkaloids of Ipecacuanha, the latter yield somewhat similar colour reactions to those of Morphine, and a further reference is made to these reactions under Emetine, Cephaeline, and P-ychotrine.

DIACETYL-MORPHINE Heroin , $\text{C}_{17}\text{H}_{17}\text{NO}$ ($\text{C}_2\text{H}_3\text{O}_2$)₂, eq 366.45—A fine white odourless crystalline powder, possessing a feeble bitter taste. Soluble 1 in 900 in Water, 1 in 40 of Alcohol (90 p.c.), readily in diluted acids.

Official in Austr (*Morphinum diaceticum*), and (Russ *Heroinum*).

Tests—The Acetic Ester of Morphine melts at 169° to 172° C (336.2° to 341.6° F). It is readily and completely soluble in Chloroform, but practically insoluble in Water. It may be distinguished from Morphine by yellow coloration when mixed with a few drops of a solution of Potassium containing a little Ferric Chloride T.S., and by its failure to set from Iodic Acid with Sulphuric Acid containing a trace of Nitric Acid it yields a yellowish-red coloration in the cold, and, on warming, a blood-red coloration, with Nitric Acid it yields at first a yellow coloration, and, on warming, a red. When warmed with a few c.c. of Alcohol (90 p.c.) it evolves a characteristic odor.

It is alkaline in reaction towards the usual indicators of neutrality, and combines with diluted mineral acids to form salts which are neutral in reaction towards the usual indicators. It may be determined by direct titration with Normal Vol. of Potassium Permanganate Solution, using Methyl Orange or Iodeosin Solution as indicator or officially 1 c.c. of Normal Volumetric Sulphuric Acid Solution represents 0.36645 gramme of Diacetyl-morphine. It should dissolve in concentrated Sulphuric Acid without colour, indicating the absence of foreign organic impurities, the presence of Morphine may be ascertained by the reaction given above. When ignited with free access of air it should leave no weighable residue.

DIACETYL-MORPHINE HYDROCHLORIDE *Heroin Hydrochloride*, $\text{C}_{17}\text{H}_{17}\text{NO}$ ($\text{C}_2\text{H}_3\text{O}_2$)₂.HCl, eq 402.64—A white, odourless, crystalline powder.

Solubility—1 in 2 of Water, 1 in 11 Alcohol (90 p.c.), insoluble in Ether.

The solubility of Heroin Hydrochloride was carefully determined in the author's laboratory, and the above figures represent the results of numerous determinations. Continental samples of the salt had a solubility of 1 in 2 of Water, which is that also given by the manufacturer.

Dott states (*C D '05*, 1 489, *P J '05*, 1 440) that the Hydrochloride appears to contain 2H₂O, one molecule of which is lost under 100° C (212° F), the remaining portion at 120° C (248° F). Samples of Continental make examined in the author's laboratory showed a loss at 100° C (212° F) of 1.2 p.c., at 120° C 1.8 p.c.

Medicinal Properties—Introduced as a substitute for Morphine, it being stated to possess the advantages of not causing constipation and of being active in much smaller doses. Has been found useful in acute and chronic bronchitis, bronchial asthma, the cough of phthisis, in acute pneumonia and in pertussis.

Result of five years' experience in the use of this drug in simple bronchitis, bronchitis with measles, the bronchitis of influenza, chronic catarrhal bronchitis, phthisis and pneumonia its effects as a cough relieving agent were prompt and definite, and in the case of almost incessant cough or severe paroxysms during the night its good effects were especially noticeable, whilst in phthisis its use was followed by most satisfactory results. In chronic bronchial catarrh it seems to have a positive curative value. In measles, when the bronchial irritation was prominent, nothing else was found as serviceable.—*J. '00*, 1 180.

Has been recommended in the treatment of the morphine habit, but its use is deprecated on the ground that the craving following its use is infinitely more unmanageable than is that of Morphine.—*B M J E '01*, 11 24, *J. '01*, 11 263, *B M J E '07*, 1 87.

Dose— $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.0027 to 0.01 gramme.

It is advisable to commence with the smaller dose, as some persons are easily affected by it, and repeated doses of $\frac{1}{16}$ and $\frac{1}{8}$ grain have produced toxic symptoms.

Foreign Pharmacopœias—Official in Jap, Swiss (*Morphinum diacetylicum hydrochloricum*), Russ (*Heronium hydrochloricum*), Russ, maximum single dose, 0.01 gramme, maximum daily dose, 0.02 gramme, Swiss, maximum single, 0.005, maximum daily, 0.15, Jap, maximum single, 0.01, maximum daily, 0.03.

Tests—Diacetyl morphine Hydrochloride has a m.p. of 116° to 117° C (240° to 242.6° F). It dissolves readily in Water, yielding solutions which are neutral in reaction towards Litmus, and from which the free base is liberated, on the addition of Ammonia Solution or Sodium Bicarbonate Solution the separated base should possess the m.p. and respond to the tests given under Heroin. The anhydrous Hydrochloride should contain theoretically 91 p.c. of Diacetyl-morphine, and the percentage of the latter may be determined by titrating a weighed quantity of the salt with Normal or Tenth normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, and adding sufficient Ether to retain the liberated alkaloid in solution. 1 c.c. of Normal Volumetric Sodium Hydroxide Solution represents 0.3645 gramme of Diacetyl morphine or 0.40264 gramme of anhydrous Diacetyl morphine Hydrochloride. Dott (*C D '05*, 1 489) has proposed the following test—Dissolve a weighed quantity of 0.4 of a gramme of the salt in 4 c.c. of Water, add 0.1 of a gramme of Sodium Bicarbonate, allowing it to stand for several hours, a precipitate should be yielded which, when collected on a small filter, washed with 6 c.c. of Water and dried in a water-bath, should weigh 0.3 of a gramme. The majority of the samples examined in the author's laboratory, when assayed according to this test, showed a higher yield of residue than 0.3 gramme. The Hydrochloride should be free from the impurities mentioned under Heroin.

ELIXIR HEROIN CUM TERPENE—Heroin, $\frac{1}{2}$ grain, Terpene Hydrate, 8 grains, Alcohol (90 p.c.), 6 fl. drms, Syrup of Virginian Prune Bark, 3 fl. drms, Glycerin, 3 fl. drms. Dose— $\frac{1}{2}$ to 2 drms = 1.8 to 7.1 c.c. (Each fl. drms contains $\frac{1}{4}$ grain Heroin and $\frac{3}{8}$ grain Terpene Hydrate).—*Bournemouth Formulary*.

Elixir Acetomorphinæ et Terpinæ—Acetomorphine Hydrochloride, 0.10, Terpin Hydrate, 1, Alcohol, 45, Glycerin, 22.50, Syrup of Wild Cherry, q.s. to produce 100.—*B P C*.

The *B P C Supplement* has changed the quantity of Alcohol to 50, and the quantity of Glycerin to 25

GLYCERINUM HEROINÆ COMPOSITUM *Syn* Glycaphorm
—Heroin Hydrochloride, 10 grains, Chloroform, 20 minims, Syrup of Roses, 10 fl oz, Distilled Water, 2 fl oz, Alcohol, 40 minims, Glycerin, *q s* to make 20 fl oz

Dissolve the Heroin in the Water and add the syrup, gradually shaking after each addition. Dissolve the Chloroform in the Alcohol, add the syrup, then add Glycerin to 20 fl oz—*Pharm Form*

Glycerinum Acetomorphinæ—Acetomorphine Hydrochloride, 0.05, Chloroform, 0.20, Alcohol (90 p c), 0.40, Syrup of Roses, 50, Distilled Water, 10, Glycerin, *q s* to make 100—*B P C*

Glycerinum Heroini Compositum—Heroin, 20 grains, Ammonium Hypophosphite, 640 grains, Fluid Extract of Hyoscyamus, 320 minims, Fluid Extract of White Pine, 2½ fl oz, Soluble Tincture of Tolu, 2 fl oz, Glycerin, 10 fl oz, Syrup of Wild Cherry Bark, 6 fl oz, Cinnamon Water, *q s* to make 40 fl oz—*Canadian Formulary* 1908

LINCTUS HEROIN—Heroin Hydrochloride, 2 grains, Tincture of Hyoscyamus, 4 fl drms, Spirit of Chloroform, 4 fl drms, Syrup of Tolu, 1 fl oz, Syrup of Wild Cherry Bark, 1 fl oz, Glycerin, *q s* to produce 6 fl oz

Mix (Each fl drms contains ⅓ grain of Heroin Hydrochloride) Dose— $\frac{1}{2}$ to 2 fl drms = 1.8 to 7.1 cc—*Bournemouth Formulary*

Linctus Acetomorphinæ—Acetomorphine Hydrochloride, 0.10, Tincture of Hyoscyamus, 7.50, Spirit of Chloroform, 7.50, Syrup of Balsam of Tolu, 15, Syrup of Wild Cherry, 15, Glycerin, *q s* to produce 100—*B P C*

PASTILLI HEROIN—Heroin Hydrochloride, 1 grain, Ammoniated Glycyrrhizin, 32 grains, Fluid Extract of White Pine Oil, 1 fl oz, Glycerin, sufficient to make 32 pastilles, each pastille contains ⅓ grain of Heroin Hydrochloride—*Bournemouth Formulary*

This has been incorporated in the *B P C* as follows—**Pastillus Acetomorphinæ Compositus**—Acetomorphine Hydrochloride, ⅓ grain, Ammoniated Glycyrrhizin, 32 grains, Pine Oil, ⅓ minim—*B P C*

BENZYL-MORPHINE HYDROCHLORIDE Peronine, $C_{17}H_{19}NO$, C_6H_5HCl , eq 408.61—It may be prepared by the action of Benzyl Chloride on Morphine

A bitter, odourless, white, micro-crystalline powder, soluble 1 in 200 of Water, 1 in 160 of Alcohol (90 p c), insoluble in Chloroform and Ether

Medicinal Properties—Narcotic and sedative, introduced as a substitute for Morphine and Codeine in the irritative cough of phthisis and chronic bronchitis—*B M J E* '98, ii 43, *L* 99, ii 139

An exhaustive report on Benzyl-morphine by Stockman and Dott, the remarks found under 'Peronine' are also applicable to the Benzyl compound—*L* '90, ii 1

Instillation of 1 to 2 p c solution into conjunctival sac induces anaesthesia of the cornea—*B M J E* '99, ii 71

Dose— $\frac{1}{2}$ to ⅓ grain = 0.008 to 0.032 grammes

Tests—Benzyl-morphine Hydrochloride, when strongly heated, evolves a strong aromatic odour, the salt dissolves in Water, forming a solution neutral in reaction to Litmus, from which Ammonium, Potassium, or Sodium Hydroxide Solution precipitates the free base. It is precipitated by the usual alkaloidal reagents, e.g., Potassium Mercuric Iodide (Mayer's) Solution, Iodo-Potassium Iodide (Wagner's) Solution, Picric Acid, etc. With Sulphuric Acid containing a trace of Nitric Acid it yields a dark brownish-red coloration. It is distinguished from Morphine by yielding no blue coloration with Potassium Ferricyanide Solution containing a few drops of Ferric Chloride T.S., and by its failure to liberate Iodine from Iodic Acid. It contains theoretically 91.1 p c of Benzyl-morphine, which may be determined by titration in a similar manner to that described under Diacetylmorphine Hydrochloride 1 cc of Volumetric Potassium

Hydroxide Solution represents 0.10861 gramme of anhydrous Benzyl morphine Hydrochloride, $C_{17}H_{19}NO_2(C_6H_5)HCl$

MONO-ETHYL - MORPHINE HYDROCHLORIDE Dionine, $C_{17}H_{19}NO_2 \cdot C_2H_5HCl \cdot H_2O$, eq 364.94—A bitter, odourless, white, micro-crystalline powder

Solubility—1 in 7 of Water, 1 in 5 of Alcohol (90 p c), insoluble in Ether

The above figures were carefully determined in the author's laboratory, and have been confirmed on repetition. The solubility of the salt in Alcohol (90 p c) varies greatly with the temperature, slight variations producing an appreciable effect, the salt which had a solubility in Alcohol (90 p c) at 15° C (59° F) of 1 in 5, dissolved readily 1 in 1 of warm Alcohol (90 p c). The figures for the solubility given by Dott (*C D '05*, i 489) are 1 in 14 of Water, 1 in 29 of Alcohol (90 p c)

Medicinal Properties—Analgesic, not a local anæsthetic

Its regular and systematic use in the form of a 5 to 10 p c solution has been recommended (*B M J '04*, ii 1303) in interstitial keratitis. A 5 p c solution was found (*L '05*, ii 835) very beneficial in corneal opacities from recent keratitis. It is one of the most valuable agents we possess (*L '06*, ii 15, *M T '07*, i 53) for the relief of deep seated ocular pain, *e g*, in glaucoma, iritis, scleritis, etc

As an analgesic (*B M J '06*, i 1098) it is used chiefly in the form of a 5 p c aqueous solution or made up with Vaseline as an ointment in similar strength. Not only an ocular anæsthetic, but a powerful ocular analgesic. It neither dilates the pupil nor increases the tension of the eye. It is advisable to begin with a 2 p c solution and gradually increase the strength so that in the course of a few days 5 p c may be used. Drops of a weak solution (1 or 2 p c) have given considerable relief in those cases, chiefly of neurotic and neurasthenic patients, where no disease or abnormality may be discoverable on careful examination, and yet the patient constantly complains of the feeling of soreness. No other method for the clearing up of corneal opacities can be compared with the results of the use of this drug. In the treatment of all forms of keratitis it may be used in 1 or 2 p c solution combined with Atropine. When the inflammatory symptoms have subsided, the Atropine is stopped, but the Dionine is continued alone for a considerable time. For the clearing up of corneal opacities, an ointment containing 4 grains to the oz may be used to commence with, gradually increasing the strength to 12 grains to the oz.

As an ocular analgesic in 5 p c aqueous solution, or as an ointment of similar strength—*B M J '04*, i 1009

Has been found useful in relieving the cough of phthisis and bronchitis—*B M J E '99*, i 36, '01, ii 68, '02, i 60, *P J '01*, ii 645

In the treatment of the drug habit, $\frac{1}{4}$ to $\frac{1}{2}$ grain doses—*B M J E '99*, i 83

Dose— $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.021 to 0.032 gramme, dissolved in Water, or in the form of a syrup

Official in Swiss Swiss, maximum single dose, 0.05 gramme, maximum daily dose, 0.15 gramme

Tests—Mono-ethyl morphine Hydrochloride melts at about 124° C (255° F). It dissolves in Water, forming a solution which is neutral in reaction towards Litmus and which yields precipitates with the usual alkaloidal agents, *e g*, Potassium mercuric Iodide (Mayer's) Solution, Iodo potassium Iodide (Wagner's) Solution, etc

Sulphuric Acid dissolves Dionine, forming a clear colourless solution, and evolves simultaneously Hydrochloric Acid gas, the addition of 1 or 2 drops of Ferric Chloride T.S. produces on warming a violet coloration changing to a deep blue, which upon the further addition of 1 or 2 drops of Nitric Acid assumes a deep red coloration. With Sulphuric Acid Solution containing 0.5 p c of Ammonium Molybdate it yields a similar violet coloration to that of Morphine. If 0.5 of a gramme of Dionine be dissolved in 5 c c of Water it yields with Ammonia Solution (sp gr 0.910) a copious white precipitate which redissolves on the addition of about 5 c c of the Ammonia Solution, but the free base again separates in the form of crystals in a short time, Codeine under similar

circumstances yields a precipitate which is permanently dissolved on the addition of 1 c.c. of the Ammonia Solution.

Ethyl-morphine may be distinguished from other alkaloids by yielding no immediate blue coloration or precipitate with a solution of Ferric chloride containing a trace of Ferric Chloride T.S. The anhydrous salt contains theoretically 85 p.c. of Ethyl-morphine, it is neutral in reaction towards Phenolphthalein and may be titrated in a similar manner to Diacetyl-morphine Hydrochloride. 1 c.c. of Normal Volumetric Sodium Hydroxide Solution represents 0.06494 gramme of anhydrous Mono-ethyl-morphine Hydrochloride. It should be free from the impurities mentioned under Diacetyl-morphine Hydrochloride.

MORPHINÆ ACETAS.

MORPHINE ACETATE

$C_{17}H_{19}NO_3$, $C_2H_4O_2$, $3H_2O$, eq 396.27

FR, ACÉTATE DE MORPHINE, GER, MORPHINACETAT, ITAL, ACETATO DI MORFINA, SPAN, ACETATO MORFICO

A light, white, crystalline powder, possessing a faint acetous odour and a bitter taste. It gradually loses Acetic Acid when exposed to the air, and should therefore be kept in well-stoppered glass bottles of a dark amber tint and exposed as little as possible to the atmosphere.

Solubility—Theoretically 1 in $2\frac{1}{2}$ of Water, but most samples will require the addition of Acid, 1 in 100 of Alcohol (90 p.c.), 1 in 5 of Glycerin.

It has been stated (*C.D.* '05, 1 282) that the solubility for Morphine Acetate would be better described as 1 in 3 of Water.

Medicinal Properties—See Morphinæ Hydrochloridum.

The *Injectio Morphinæ Hypodermica* formerly (*B.P.* 1885) contained one grain of Morphine Acetate in ten minims, now (*B.P.* 1898) it contains one grain of Morphine lactate in twenty-two minims.

Recommended in diabetes—*Pr.* xxxviii 20, *B.M.J.* '89, 1 118.

Dose— $\frac{1}{2}$ to $\frac{1}{4}$ grain = 0.008 to 0.032 gramme.

Prescribing Notes—As it is very liable to lose the acid, dry the salt without a slight loss of Acetic Acid, the commercial Acetate generally requires a little added Acetic Acid to make a clear solution. Aqueous solutions have a strong tendency to deposit a basic Morphine Acetate, and to become acid.

Incompatibles—Alkalis and alkaline earths, astringent vegetable infusions and decoctions.

Official Preparation—Liquor Morphinæ Acetatis.

Not Official—*Injectio Morphinæ et Atropinæ*.

Antidotes—See Morphine Hydrochloride.

Foreign Pharmacopœias—Official in Belg, Mex, Port and U.S. Not in the others.

Tests.—Morphine Acetate when heated loses Water and Acetic Acid and melts, according to the *U.S.P.*, at about 200° C. (392° F.).

It affords upon the addition of Ammonia Solution, in slight excess, a white precipitate rapidly becoming crystalline, and if this precipitate is separated, washed with a little cold Morphinated Water and dried, it responds to the tests described under Morphine. A small quantity of the salt warmed with a little Sulphuric Acid and 1 or 2 cc of Alcohol (90 p c) yields a characteristic odour of Ethyl Acetate. The test for Acetates with Ferric Chloride Test solution cannot be employed unless the alkaloid is first removed, owing to the reaction of the Morphine which produces a greenish blue destroyed by acids or by heat. When warmed with Sulphuric Acid it evolves an odour of Hydrogen Acetate. It is officially required to yield 71 p c of anhydrous Morphine equivalent to 75.4 p c of hydrated Morphine and to 99.4 p c of a salt of the pharmacopoeial formula, as gravimetrically determined by dissolving 2 grammes of the salt in a mixture of 6 cc of warm Morphinated Water and 0.1 cc of Acetic Acid and precipitating this solution with Ammonia Solution in slight excess, the precipitate being washed with a little cold Morphinated Water, dried first by pressure between sheets of bibulous paper, and then at a temperature of 55° to 60° C (131° to 140° F), and eventually at a temperature of 110° C (230° F), the crystals should weigh 1.42 grammes. The *B P* states that in the event of the salt yielding a larger proportion of Morphine than this, before use it should be recrystallised from hot Water acidulated with Acetic Acid. It should be free from the impurities mentioned under Morphine, when ignited with free access of air the salt should leave no weighable residue, indicating the absence of mineral impurity.

Preparation

LIQUOR MORPHINÆ ACETATIS. SOLUTION OF MORPHINE ACETATE

Morphine Acetate, 17½ grains, Diluted Acetic Acid, 38 minims, Alcohol (90 p c), 1 fl oz, Distilled Water, *qs* to yield 4 fl oz (1 in 100)

Dose — 10 to 60 minims = 0.6 to 3.6 cc

11 minims contain ⅓ grain

Not Official

INJECTIO MORPHINÆ ET ATROPINÆ HYPODERMICA — Morphine Acetate, 10 grains, Atropine Sulphate, ⅓ grain, Water, 120 minims, dissolve ⅓ grain of Morphine Acetate and ⅓ grain of Atropine Sulphate in every 6 minims

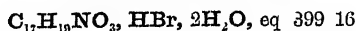
Dose — 1 to 6 minims for each injection = 0.06 to 0.36 gramme

Atropine combined with Morphine increases its analgesic and hypnotic effects, whilst it lessens the tendency to sickness, dyspepsia, depression and constipation.

The *B P C* solution contains 6 p c of Morphine Sulphate and 0.12 p c of Atropine Sulphate

Not Official
MORPHINÆ HYDROBROMIDUM.

MORPHINE HYDROBROMIDE



Forms long, colourless needles, soluble 1 in 25 Water, 1 in 50 of Alcohol (90 p c). It is employed for similar purposes to the Hydrochloride.

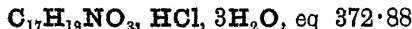
Dose — $\frac{1}{2}$ to $\frac{1}{2}$ grain = 0.008 to 0.032 gramme

Tests — Morphine Hydrobromide loses of 100° to 110° C. (212° to 230° F) two molecules of Water of crystallization, equivalent to 8.9 p c. It dissolves in Water, yielding a solution which is neutral or but faintly acid to Litmus paper, on the addition of a slight excess of Ammonia Solution it yields a precipitate rapidly becoming crystalline, which, when separated, washed and dried, answers the tests characteristic of Morphine given under that heading. The aqueous solution yields with Silver Nitrate Solution a yellowish curdy precipitate which, when separated and washed, dissolves readily in Potassium Cyanide Solution, but which is practically insoluble in Ammonia Solution and insoluble in Nitric Acid. It contains theoretically 70.9 p c of anhydrous Morphine, equivalent to 75.4 p c of hydrated Morphine, which may be gravimetrically determined by methods similar to those described under Morphine Acetate, *U. S. P.* Titrate. It should be free from the impurities mentioned. The salt, when ignited with free access of air, should leave no weighable residue.

MORPHINÆ HYDROCHLORIDUM.

MORPHINE HYDROCHLORIDE

HYDROCHLORATE OF MORPHINE — *B. P.* '85



FR, CHLORHYDRATE DE MORPHINE, GER, MORPHINHYDROCHLORID, ITAL, CLORIDRATO DI MORFINA, SPAN, CLORURO MORFICO

White, odourless, lustrous, silky needles or a white micro-crystalline powder, possessing a bitter taste.

Solubility — 1 in 24 of Water, about 1 in 72 of Alcohol (90 p c), 1 in 8 of Glycerin, insoluble in Ether.

The above figure for Morphine Hydrochloride in Water has been confirmed on repetition the figure for Alcohol (90 p c) has been altered from 1 in 50 to 1 in 72, the former figure represented the solubility in Rectified Spirit of the *B. P.* 1885, the increased strength of the Alcohol official in *B. P.* 1898 causing difference in solubility. The Report of the Committee of Reference in Pharmacy recommends the solubility of 1 in 25 of Water, and states that 1 gramme dissolves in 69 c c of Alcohol (90 p c). It should be kept in well-stoppered bottles of a dark amber tint.

Medicinal Properties — Morphine possesses in a marked degree the analgesic and hypnotic effects of Opium. It has the advantage over Opium of being less apt to disturb digestion and cause constipation, more particularly when given by hypodermic injection; it is also less likely to cause headache and nausea. It is more readily absorbed and acts quicker, it is better adapted for hypodermic

injection and for suppositories. Opium, however, is better for relieving pain in the alimentary tract, as in gastric ulcer and abdominal pain, it is also more useful as a diaphoretic and in diabetes. It lessens the secretions, diminishes diarrhoea, and produces constipation. Children are very susceptible to the action of Morphine.

Valuable papers on its use in epidemic diseases, *J.* '98, ii 1393, and on Opium in acute and chronic disease, *Ph.* '07, i 625.

Given immediately before the general anæsthetic or before the patient leaves the table, prevents post anæsthetic vomiting.—*J.* '08, i 292.

A hypodermic injection given before the administration of a general anæsthetic decreases the patient's susceptibility to shock during the operation.—*L.* '05, i 853.

Amongst the various references to Morphine or Morphine salts, $\frac{1}{4}$ to $\frac{1}{2}$ grain has been recommended hypodermically (*B M J* '04, ii 1635, 1783) in the treatment of hæmoptysis. An exception to the exhibition of Morphine (*B M J* '05, i 68) is found when the bleeding is so profuse as to flood the air passages, and threatens to suffocate the patient.

Morphinomania treated successfully by Atropine and Strychnine.—*B M J* '07, i 1173.

In the treatment of puerperal eclampsia, injection of $\frac{1}{2}$ to 1 grain.—*B M J* '01, ii 810, '02, i 71, 509, '05, ii 718, 719, 749, *J.* '01, i 1823, *T G* '01, 622.

Dose— $\frac{1}{8}$ to $\frac{1}{2}$ grain = 0.008 to 0.032 gramme.

Ph. Ger. maximum single dose, 0.03 gramme, maximum daily dose, 0.1 gramme.

Prescribing Notes—*The salts of Morphine are all readily soluble in Water, the Acetate being the most soluble, but it is apt to deposit a basic salt, the Tartrate and Lactate are next, being about 1 in 10, the Sulphate, Hydrobromide and Hydrochloride are the least soluble, requiring rather more than 20 of Water to 1 of the salt.*

Incompatibles—Alkalis and alkaline earths, astringent vegetable infusions and decoctions, Ferric Chloride.

Official Preparations—Liquor Morphinæ Hydrochloridi, Suppositoria Morphinæ, Trochiscus Morphinæ and Trochiscus Morphinæ et Ipecacuanhæ. Contained in Tinctura Chloroformi et Morphinæ Composita.

Not Official—Insufflatio Morphinæ, Linctus Morphinæ (*Squire*), Linctus Morphinæ Acidus (*Squire*).

Antidotes—If taken by the mouth, induce vomiting, and wash out the stomach. Keep the patient walking about, and rouse him in every way. Ammonia or Spirit of Sal Volatile to the nose, inject a pint of strong Coffee into the bowel. Hypodermic injection of Atropine Sulphate $\frac{1}{20}$ grain, repeating in a quarter of an hour if necessary. Tincture of Belladonna, Amyl Nitrite inhalation, artificial respiration.—*Munell* $\frac{3}{4}$ grain Strychnine acts as an antidote to $\frac{1}{2}$ grain Morphine.—*L.* '71, ii 840, 907. Potassium Permanganate is used to wash out the stomach, a solution of 120 minims of Liq. Pot. Permang. in a pint of Water is suitable. If quantity of Opium or Morphine taken is unknown, 8 to 10 grains Potassium Permanganate in from 4 to 8 fl. oz. of Water should be administered at once. The solution may be acidulated with Acid Sulphuricum Dilutum with advantage.—*B M J* '95, i 1369, '95, ii 55, 76, '96, i 1194, *T G* '98, 97. Picric acid, $\frac{1}{10}$ grain.—*L.* '89, i 497. $\frac{1}{2}$ grain doses of Cocaine at intervals of half an hour until consciousness returns and breathing is normal, as an antidote.—*M P* '02, i 147, *P J* '02, i 114.

Case of poisoning by 8 grains of Morphine Sulphate (in three hypodermic injections), treated by artificial respiration, subcutaneous injection of 30 oz. of normal saline solution, and drinking a diluted solution of Potassium Permanganate.—*L.* '02, i 1317.

Value of Oxygen in poisoning by Morphine.—*L.* '98, ii 545.

Foreign Pharmacopœias - Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung Ital, Jap Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests —Morphine Hydrochloride when heated to 100°C (212°F) loses its Water of Crystallization equivalent to 14.4 p c, and when still more strongly heated chars without melting. The German Pharmacopœia states that it loses 14.4 p c of its weight at 100°C (212°F), and that the anhydrous Morphin Hydrochloride should be of a pure white or only of a pale yellow colour, when heated to 250°C (482°F) it changes to a brown colour.

It dissolves in Water, yielding a solution neutral in reaction towards Litmus paper, when the solution is treated with a slight excess of Ammonia Solution it yields a white precipitate rapidly becoming crystalline which, when separated and washed with Morphinated Water and carefully dried, answers tests characteristic of Morphin given under that heading. The aqueous solution acidified with Nitric Acid yields on the addition of Silver Nitrate Solution a white curdy precipitate, which when separated dissolves readily in Ammonia Solution and in solution of Potassium Cyanide, but is insoluble in Nitric Acid. When warmed with Sulphuric Acid it evolves Hydrochloric Acid gas. It is officially required to contain 75.5 p c of anhydrous Morphin, equivalent to 80.2 p c of hydrated Morphin and to 99.5 p c of pure Morphin Hydrochloride of the pharmacopœial formula as determined by dissolving 2 grammes of the salt in 250 c c of warm Morphinated Water, the Morphin with the smallest possible excess of Ammonia Solution, filtering, washing the crystalline precipitate with a little cold Morphinated Water, and drying first between folds of bibulous paper, then at a temperature of 55 to 60°C (131° to 140°F) and finally at a temperature of 110°C (230°F), the crystals should weigh 1.51 grammes.

The Report of the Committee of Reference in Pharmacy recommends that the 2 grammes of the salt employed in the above quantitative estimation should be dissolved in 50 c c instead of 250 c c of warm Morphinated Water, and the precipitated Morphin should weigh 1.5 to 1.51 grammes. Morphin Hydrochloride should be free from the impurities mentioned under Morphin. The *U.S.P.* states that on the addition of Potassium Carbonate T.S. to a 1 in 30 aqueous solution of the salt a white precipitate should be formed, which should be soluble without colour in Chloroform indicating the absence of Apomorphine. The Benzol test is given under Morphin.

Potassium Carbonate—5 c c of an aqueous solution of the salt (1-30) should give immediately, or after a few seconds, with 1 drop of Potassium Carbonate T.S., a pure white crystalline precipitate. On exposure to the air undergoes no coloration. When agitated with Chloroform no red colour should be developed, *P.G.*, the white precipitate formed should dissolve in Chloroform without colour, *U.S.P.*

Ether—If the solution of this precipitate in Sodium Hydroxide T.S. be agitated with an equal volume of Ether, the separated ethereal layer when evaporated to dryness should not leave a weighable residue, *P.G.*

Potassium Hydroxide—An aqueous solution of the salt yields a white precipitate with Potassium Hydroxide T.S., readily soluble in excess, *B.P.*

Preparations

LIQUOR MORPHINÆ HYDROCHLORIDI SOLUTION OF MORPHINE HYDROCHLORIDE *B P Syn*—SOLUTION OF HYDROCHLORATE OF MORPHINE

Morphine Hydrochloride, 17½ grains, Diluted Hydrochloric Acid 38 minims, Alcohol (90 p c), 1 fl oz, Distilled Water, q s to yield 4 fl oz (1 in 100)

Dose—10 to 60 minims = 0.6 to 3.6 c c

11 minims contain $\frac{1}{10}$ grain of Morphine Hydrochloride

Foreign Pharmacopœias—Official in *Fr* Soluto de Morphine (Chlorhydrate) pour Injection Hypodermique, 1 in 50, *Port* (Soluto de Chlorhydrate de Morphina), 1 in 20, for hypodermic injection Not in the others

SUPPOSITORIA MORPHINÆ MORPHINE SUPPOSITORIES

$\frac{1}{4}$ grain of Morphine Hydrochloride in each, with Oil of Theobroma

Half the strength of *B P* '85

TINCTURA CHLOROFORMI ET MORPHINÆ COMPOSITA

The formula is given under Chloroform The proportion of Morphine has been much increased, and is now more than four times what it was in *B P* '85 10 minims now contain $\frac{1}{2}$ minim of Chloroform, $\frac{1}{2}$ Minim of Diluted Hydrocyanic Acid, $\frac{1}{10}$ grain of Morphine Hydrochloride, and 1 minim of Tincture of Indian Hemp

TROCHISCUS MORPHINÆ MORPHINE LOZENGE

$\frac{1}{10}$ grain of Morphine Hydrochloride in each lozenge, with Tolu Basis

Dose—1 to 6 lozenges One occasionally for cough

Official in Jap (Pastilli Morphini Hydrochlorici), 0.005 gramme in each pastille

TROCHISCUS MORPHINÆ ET IPECACUANHÆ MORPHINE AND IPECACUANHA LOZENGE

$\frac{1}{10}$ grain of Morphine Hydrochloride, and $\frac{1}{12}$ grain of powdered Ipecacuanha Root in each, with Tolu Basis

Dose—1 to 6 lozenges One occasionally for cough

Foreign Pharmacopœias—Official in *U S* contains $\frac{1}{10}$ grain of Morphine Sulphate, and $\frac{1}{12}$ grain of Ipecacuanha in each, *Swiss* (Pastilli Ipecacuanhæ cum Opio), contains about 0.002 gramme = $\frac{1}{12}$ grain of each, *Ipecac* and *Opium Jap* (Pastilli Opii et Ipecacuanhæ) Each pastille contains 0.025 gramme = about $\frac{1}{10}$ grain of each, *Opium* and *Ipecacuanha*

Not Official

INSUFFLATIO MORPHINÆ—Morphine Hydrochloride, $\frac{1}{2}$ grain, Bis-muth Oxychloride, 1 grain, Starch, $\frac{1}{2}$ grain—*City Chest*

The Morphine Insufflations of *Royal Chest* are dilutions with Milk Sugar, and those of *Throat and Great Nostril*, with dried Starch

LINCTUS MORPHINÆ (*Squire*)—Solution of Morphine Hydrochloride, 3 minims, Spirit of Chloroform, 3 minims, Glycerin, 30 minims, Syrup, to 1 fl drim

LINCTUS MORPHINÆ ACIDUS (*Squire*)—Solution of Morphine Hydrochloride, 3 minims, Spirit of Chloroform, 3 minims, Lemon Juice, 15 minims, Glycerin, to 1 drim

Both the above are very palatable

This has been incorporated in the *B, P C* under title **Linctus Sedativus**
Syn—Linctus Morphinæ Acidus

Not Official

MORPHINÆ LACTAS.

Morphine Lactate, $C_{17}H_{19}NO_3 \cdot C_3H_5O_3$, eq 372.42, occurs in colourless, prismatic crystals

Solubility —1 in 8 of Water, 1 in 93 of Alcohol (90 p c)

Dose — $\frac{1}{2}$ to $\frac{1}{2}$ grain = 0.008 to 0.032 gramme

Not Official

MORPHINÆ SULPHAS

Morphine Sulphate $(C_{17}H_{19}NO_3)_2 \cdot H_2SO_4 \cdot 5H_2O$, eq 752.84, occurs in colourless acicular crystals

Solubility —1 in 21 of Water, freely in hot Water, 1 in 700 of Alcohol (90 p c)

Dose — $\frac{1}{2}$ to $\frac{1}{2}$ grain = 0.008 to 0.032 gramme

Foreign Pharmacopœias —Official in Jap, Mex, Norw, Port, Span and U S Not in the others

Tests —Morphine Sulphate loses 3 molecules of Water of crystallisation, equivalent to 7.12 p c at 100° C (212° F). When strongly heated (about 250° C (482° F)) it changes to a brown colour and finally chars without melting. It dissolves in Water, yielding a clear solution which is neutral in reaction towards Litmus paper; this solution affords with Ammonia Solution a white crystalline precipitate, which should answer the tests characteristic of Morphine given under that substance. It should afford with Barium Chloride T.S. a white precipitate insoluble in Hydrochloric Acid. It contains theoretically 75.2 p c of anhydrous Morphine, and may be gravimetrically determined by a similar process to that given for Morphine Acetate, Hydrochloride, or Tartrate.

LIQUOR MORPHINÆ SULPHATIS —Sulphate of Morphine, 1, Rectified Spirit 25, Distilled Water, *q s* to produce 100 —*B P* 1885

This has been incorporated in the *B P C*

PULVIS MORPHINÆ COMPOSITUS Morphine Sulphate 15, Camphor, 32, Glycyrrhiza in No. 80 powder, 33, Precipitated Calcium Carbonate, 33.5, Alcohol, *q s*. Rub the Morphine Sulphate with the Precipitated Calcium Carbonate, added in portions of about 5 each, until it is thoroughly mixed, then rub the Camphor with a little Alcohol and mix intimately with the Glycyrrhiza and the other powder. Finally, pass the powder through a No. 40 sieve, pulverise the residue if any should be left on the sieve, add to the sifted powder and mix thoroughly. Transfer it to well-stoppered bottles. Average Dose, $7\frac{1}{2}$ grains = 0.5 gramme —*U S P*

This has been incorporated in the *B P C*

PILULÆ ATROPINÆ ET MORPHINÆ See p 204

MORPHINÆ TARTRAS.

MORPHINE TARTRATE.

$(C_{17}H_{19}NO_3)_2 \cdot C_4H_6O_6 \cdot 3H_2O$, eq 768.66

FR, TARTRATE DE MORPHINE, GER, TARTRAT VON MORPHIN, ITAL, TARTRATO DI MORFINA, SPAN, MORFINO TARTRATO

Colourless, acicular crystals, or as a fine, white, crystalline powder

It may be prepared by the combination of Morphine with its molecular equivalent of Tartaric Acid.

Solubility—1 in 10 of Water, sparingly in Alcohol (90 p c)

The solubility of this salt is sometimes affected by the presence of a small quantity of the Acid Tartrate, the latter being much less soluble than the official salt

Dose— $\frac{1}{8}$ to $\frac{1}{2}$ grain = 0.008 to 0.032 gramme

Official Preparations—*Injectio Morphine Hypodermica* and *Liquor Morphine Tartratis*

Tests—Morphine Tartrate dissolves in Water, forming a solution which is neutral in reaction towards Litmus paper, the solution yields, on the addition of Ammonia Solution in slight excess, a white precipitate rapidly becoming crystalline, which when separated yields the tests distinctive of Morphine described under that heading. The aqueous solution after the removal of the alkaloid, when mixed with Calcium Chloride Solution in excess, affords a white granular precipitate which, when separated and washed, is soluble in concentrated Potassium Hydroxide Solution, when a sufficiently concentrated solution is mixed with concentrated Potassium Acetate Solution it affords, when acidulated with Acetic Acid, a white crystalline precipitate, the precipitation being still more marked in the presence of Alcohol (90 p c), Ferrous Sulphate Solution added to a solution acidulated with Acetic Acid, followed by the addition of a few drops of Hydrogen Peroxide Solution, and finally with an excess of Potassium Hydroxide Solution, gives a purple or violet coloration. It is officially required to yield 73.5 p c of anhydrous Morphine equivalent to 78.13 p c of hydrated Morphine as gravimetrically determined by dissolving 2 grammes of the salt in 20 c c of warm Morphinated Water, precipitating the alkaloid with the slightest possible excess of Ammonia Solution, washing the crystalline precipitate with cold Morphinated Water and drying first between folds of bibulous paper, then at a temperature of 55° to 60° C (131° to 140° F), and finally at a temperature of 110° C (230° F), the crystals should weigh 1.47 grammes. It should be free from the impurities mentioned under Morphine, and when ignited with free access of air the salt should leave no weighable residue.

The salt is efflorescent at 20° C (68° F), and it should therefore be kept in well-stoppered glass bottles of a dark amber tint and kept as far as possible from contact with the air and in a cool place. It has a tendency, when kept for a lengthened period, to become converted into the Acid Tartrate, which dissolves with less facility in Water, and the presence of any appreciable proportion of such a salt is at once manifest by the increased insolubility in Water.

Preparations**INJECTIO MORPHINÆ HYPODERMICA** HYPODERMIC INJECTION OF MORPHINE

Dissolve 50 grains of Morphine Tartrate in recently boiled and cooled Distilled Water to produce 1100 minims (1 in 20)

Dose—By subcutaneous injection, 2 to 5 minims = 0.12 to 0.3 c c

The official Hypodermic Injection is now made in the Tartrate containing 1 grain in 22 minims, which is about half the strength of that of B.P. 85. On account of the great difference in strength of the various salts of Morphine, it is extremely important that they should be very plainly labelled, and used with very great care. It is also desirable that prescribers should clearly define the strength.

Tablets for making the injection are convenient and portable.

Atropine Sulphate, $\frac{1}{100}$ to $\frac{1}{1000}$ grain is added to each dose of Morphine Injection to increase its analgesic and hypnotic effects, and to lessen its depressing and constipating effects.

LIQUOR MORPHINÆ TARTRATIS. SOLUTION OF MORPHINE TARTRATE

Morphine Tartrate, $17\frac{1}{2}$ grains, Alcohol (90 p.c.), 1 fl oz, Distilled Water, *q.s.* to yield 4 fl oz (1 in 100)

Dose — 10 to 60 minims = 0.6 to 3.6 c.c.
11 minims contain $\frac{1}{10}$ grain

MORRHUÆ OLEUM.

COD-LIVER OIL

NO Syn—OLEUM JECORIS ASELLI

Fr., HUILE DE FOIE DE MORUE, GER., LIEBERTHAN, ITAL., OLIO DI FEGATO DI MERLUZZO, SPAN., ACEITE DE HIGADO DE BACALAO

A pale yellow, or yellow oily fluid, possessing a characteristic fishy odour and taste, extracted from the liver of the Cod, *Gadus Morhua*, at a temperature not exceeding 82° C (180° F), subsequently removing solid fat by filtration at about -5° C (23° F).

The alkaloids, Morrhaine and Aselline have been isolated.

Solubility.—Sparingly in Absolute Alcohol, 1 in 2 of Ether, 1 in 3 of Acetic Ether.

A solution of pure Quinine 1 fl oz at 140° F will dissolve 4 grains readily.

Medicinal Properties.—Nutritive, nervine and hæmatinic tonic. Most efficient in all forms of tubercular disease and in rickets and tertiary syphilis, useful in the chronic eczema and chronic bronchitis of children, and generally in all cases of impaired nutrition and nervous debility due to over-work, acute disease, or under-feeding. In pulmonary consumption it deservedly possesses a high reputation given in emulsion, with or without Malt Extract. It is contra-indicated in hæmoptysis, diarrhoea, and dyspepsia. It is easily assimilated, and is best given after meals, but it may produce indigestion and nausea, sometimes administered by munction, but the odour is objectionable.

It is stated (*Edin Med Journ* '05, 468) to be pre-eminently the best of the tonic remedies which have been used in the treatment of pulmonary tuberculosis. It should be taken twice a day within half an hour after meals. Best to commence with 1 drim doses and after a time to increase to 2 drim doses, the amount should never exceed $\frac{1}{2}$ oz. For children Parrish's Food or Lion Wine is a good vehicle.

Dose — 1 to 4 fl drim = 3.6 to 14.2 c.c.

Prescribing Notes.—Numerous and varied methods have been adopted for covering the taste of the Oil. It has been given floating on Orange Wine, Orange

Juice Mill, Coffee, also in a mixture of Tincture of Orange diluted mineral Acid and Syrup. A favourite form is that of the Emulsion which may be made with Gum Acacia, Tragacanth, or yolk of egg or any combination of these another and very excellent method is that given under Malt Extract, see p 270. The Oil by itself may be given in capsules containing 30 or 60 minims in each

Not Official—Cremor Morrhue Pancreaticus, Emulsion d'Huile de Foie de Morue, Emulsio Olei Morrhue, Emulsio Olei Morrhue cum Hypophosphitibus, Ovis et Vitis, Emulsio Morrhue Pancreatica, Emulsio Morrhue Pancreatica cum Extracto Malti, Emulsio Olei Morrhue Composita, Emulsium Olei Morrhue cum Hypophosphitibus, Huile de Foie de Morue Ciccosotie, Huile de Foie de Morue Phosphoree, and Morrhual

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Ger, Hung, Norw, Russ, Swed and Swiss (Ol Jecoris Aselli), Fr (Huile de Foie de Morue), Ital (Oleo di Fegato di Merluzzo), Jap (Ol Jecoris), Port (Oleo de Bacalhau), Mex and Spau (Aceite de Hígado de Bacalao), U S (Oleum Morrhue). Dutch and Swed have also Oleum Jecoris Aselli Ferratum. Dutch has also Oleum Jecoris Aselli cum Iodeto Ferroso, Swiss has also Oleum Jecoris Iodatum

Tests—Cod-Liver Oil is officially required to possess a sp gr of 0.920 to 0.930. Fifteen samples examined in the author's laboratory had a sp gr of 0.922 to 0.928, with an average of 0.925. The *USP* states 0.918 to 0.922 at 25° C (77° F), the *PG* 0.926 to 0.931. The Oil is officially required to be readily soluble in Ether and in Chloroform and slightly soluble in Alcohol (90 p c). A violet coloration is developed when a drop of Sulphuric Acid is added to a few drops of Oil on a porcelain tile. The *USP* includes a test in which Sulphuric Acid is added to a chloroformic solution of the Oil, the *PG* one in which the Acid is added to the Carbon Bisulphide Solution of Oil. The *BP* requires that a precipitate of coagulated Albumen should be formed at the junction of the two liquids when Nitric Acid is carefully poured on to the surface of some Oil contained in a test-tube. Neither the *USP* nor *PG* includes a test for Albumen with Nitric Acid. Exposure of the Oil for two hours to a temperature of 0° C (32° F) should not result in the separation of solid fat. The *PG* states that no solid fat, or only an insignificant amount, should separate on prolonged standing at 0° C (32° F). The Saponification value, the Iodine absorption and a determination of the amount of free fatty acid are important factors to be observed in judging of the purity of a specimen of Cod Oil. Cod-Liver Oil has a Saponification value of about 185, and an Iodine absorption of about 160. Fifteen samples examined in the author's laboratory had Saponification values from 181.4 to 190.4, with an average of 186.8, and Iodine absorptions from 147.3 to 165.1, with an average of 155.7. The *USP* requires a Saponification value of 175 to 185, and an Iodine absorption of not less than 140 nor more than 150, the *PG* a Saponification value of not more than 196.5 and an Iodine absorption of not less than 140 nor more than 152. The percentage of free acid varies from 0 p c in a very fine colourless Oil to 9 p c in dark-coloured samples altered by heat and long keeping, it may be determined by warming a weighed quantity of 5 or 10 grammes of the Oil with 20 c c. of Alcohol (90 p c) and titrating with Tenth-normal Volumetric Sodium Hydroxide Solution,

1 cc of the Tenth-normal solution being equivalent to 0.028011 gramme of Oleic Acid. Good specimens require from 0.0 to 1.8 cc (average 1.3 cc) of Tenth-normal Volumetric Sodium Hydroxide Solution to neutralise the free acid contained in 5 grammes of the Oil, corresponding to from 0.0 to 1.0 pc (average 0.73 pc), of free Acid calculated as Oleic Acid. Fifteen samples examined in the author's laboratory showed from 0.0 to 1.0 pc of free Oleic Acid. The *USP* and *PG* limit of free Acid is that the Oil shall at the most have only a very slightly acid reaction towards blue Litmus paper moistened with Alcohol of the respective pharmacopœial strengths. Upon the acidity of the sample also depends the presence or absence of Albumen, fine oils with little acid show an Albumen ring on being floated upon Nitric Acid, sp. gr. 1.400.

The determination of the refractive index has been suggested (*CD* '02, 1505) as a means of judging the purity of the Oil. The Norwegian Oil is stated to have a higher refractive index than Newfoundland Oil. The refractive figure of Newfoundland Oil is shown to vary between +42 to +44.45, whilst that of the Norwegian Oil varies between +44 to +48. Cod-Liver Oil produces a great increase in temperature when mixed with Sulphuric Acid (Maumené), which distinguishes it from most other oils except liver oils. For the detection of Seal Oil and other fish oils the Nitric Acid colour test may be employed. Vegetable oils may be detected by the alteration in density and the reduction in the Iodine absorption. Refined Seal Oil, which has been frequently used as an adulterant, may be detected by the determination of the Saponification value, the Iodine absorption and rise of temperature with Sulphuric Acid.

Cod-Liver Oil contains from 1 to 1.5 pc of unsaponifiable matter, which may be determined by saponifying 5 grammes of the Oil with sufficient Alcoholic Potassium Hydroxide Solution to ensure complete saponification, evaporating off the Alcohol on a water-bath, dissolving the residue in hot Water, transferring to a separator, cooling, and when cold shaking out with Ether, the Ether is evaporated off, the residue dried at 100° to 105° C (212° to 221° F) till constant in weight, when dissolved in a little Carbon Bisulphide and tested with Sulphuric Acid it should give a well-marked purple coloration.

Litmus—The Oil should show only a very slightly acid reaction to blue Litmus paper previously moistened with Alcohol, *PG* and *USP*.

Sulphuric Acid—A few drops of the Oil on a white slab give a violet coloration with a drop of strong Sulphuric Acid, *USP* and *PG*. A solution of 1 drop of Oil in 20 drops of Chloroform, shaken with 1 drop of Sulphuric Acid, acquires a violet-red tint, rapidly changing to rose-red and finally to brownish-yellow, *USP*. A solution of 1 drop of the Oil in 20 drops of Carbon Bisulphide, coloured violet-red on the addition of 1 drop of Sulphuric Acid, afterwards changing to brown, *PG*.

Nitric Acid—A mixture of 15 drops of Oil and 3 drops of fuming Nitric Acid gives a light rose coloration, changing to lemon-yellow, *PG*. If 2 or 3 drops of Nitric Acid be allowed to flow alongside of 10 to 15 drops of the Oil, contained in a watch glass, a red colour will be produced at the point of contact, when the mixture is stirred, the colour becomes bright rose-red, changing soon to lemon yellow. Seal Oil shows at first no change of colour, whilst other fish

oils when examined by this test become at first blue and afterwards brown and yellow, *U S P*

Elaidin Test—If a mixture of 1 c c of fuming Nitric Acid, 1 c c of Water and 2 c c of Cod Liver Oil be carefully shaken together, it should neither wholly nor in part solidify within 1 or 2 days, *P G*

Iodine Absorption—Dissolve about 0.3 gramme of the Oil in 10 c c of Chloroform in a 250 c c flask and add 25 c c of a mixture of equal parts of Alcoholic Iodine T S and Alcoholic Mercuric Chloride T S. Allow the mixture to stand for 4 hours protected from light, then add 20 c c of Potassium Iodide T S and dilute with 50 c c of Water. Then, on titrating with Tenth-normal Volumetric Sodium Thiosulphate Solution, the Iodine value of not less than 140 nor more than 150 should be obtained, *U S P*; about 0.5 gramme of the Oil is weighed into a stoppered bottle, dissolved in 15 c c of Chloroform, and 25 c c of Alcoholic Iodine Solution and Alcoholic Mercuric Chloride Solution added, and the mixture allowed to stand for 4 hours in a dark chamber protected from direct daylight, 1.5 gramme of Potassium Iodide and 100 c c of Water are added to the mixture, and the excess of Iodine is titrated with Tenth-normal Volumetric Sodium Thiosulphate Solution, 100 parts by weight of Cod Liver Oil shall absorb not less than 140 and not more than 152 parts of Iodine, *P G*

Saponification—If 1 gramme of the Oil be heated in a reflux condenser with 20 c c of Semi-normal Volumetric Potassium Hydroxide Solution for half an hour, and, after cooling and the addition of a few drops of Phenolphthalein T S, the mixture be titrated with Semi-normal Hydrochloric Acid Volumetric Solution, not less than 13 c c of acid should be required for decoloration, *P G*

Not Official

***CREMOR MORRHUÆ PANCREATICUS**—Stronger Glycerin of Pepsin, 4, Glycerin of Pancreatin, 4, Cod Liver Oil, 50, Decoction of Irish Moss, 27.50, Syrup of Tolu, 3, Alcohol, 3, Essential Oil of Almonds, 0.10, Distilled Water, *q s* to produce 100—*B P C*

***EMULSIO MORRHUÆ PANCREATICA**—Glycerin of Pancreatin, 2.50, Stronger Glycerin of Pepsin, 2.50, Cod Liver Oil, 40, Gluside, 0.025, Solution of Potassium Hydroxide, 1, Tragacanth, in powder, 2, Gum Acacia, in powder, 8, Oil of Cassia, 0.075, Oil of Bitter Almonds, 0.075, Distilled Water, *q s* to produce 100—*B P C*

B P C Supplement has altered the quantities as follows—Glycerin of Pancreatin, 3.5, Stronger Glycerin of Pepsin, 3.5, Cod Liver Oil, 50, Gluside, 0.033, Solution of Potassium Hydroxide, 1.25, Tragacanth, in powder, 2.5, Gum Acacia, in powder, 10, Oil of Cassia, 0.1, Oil of Bitter Almonds, 0.1, Distilled Water, *q s* to produce 100

***EMULSIO MORRHUÆ PANCREATICA CUM EXTRACTO MALTI**—Glycerin of Pancreatin, 9, Cod Liver Oil, 40, Gum Acacia, in powder, 2, Tragacanth, in powder, 0.25, Saccharated Solution of Lime, 2, Extract of Malt, *q s* to produce 100—*B P C*

B P C Supplement has altered the quantities as follows—Glycerin of Pancreatin, 10, Cod Liver Oil, 50, Gum Acacia, in powder, 2.5, Tragacanth, in powder, 0.31, Saccharated Solution of Lime, 2.5, Extract of Malt, *q s* to produce 100 by volume

EMULSIO OLEI MORRHUÆ—Cod Liver Oil, 8 fl oz, the Yolks of two Eggs, Tragacanth, in powder, 16 grains, Elixn of Gluside, 60 minims Simple Tincture of Benzoin, 60 minims, Spirit of Chloroform, $\frac{1}{2}$ fl oz, Essential Oil of Bitter Almonds, 8 minims, Distilled Water, to produce 16 fl oz. Measure 5 fl oz of the Water, place the Tragacanth in a dry mortar and triturate with a little of the Cod Liver Oil, then add the yolks of Eggs and stir briskly, adding Water as the mixture thickens. When of a suitable consistency, add the remainder of the Oil and Water alternately, with constant stirring, avoiding frothing

* These formulas closely resemble those previously published in *Annals of Formulary* (9th edition)

Put in a pint bottle, add the Elixir of Gluside, Tincture of Benzoin, Spirit of Chloroform and Oil of Almonds, previously mixed, shake well, and add Distilled Water if necessary to make the product measure 16 fl oz — *B P C Formulary* 1901, now incorporated in *B P C* as follows —

Emulsio Olei Morrhue Composita—Cod-Liver Oil, 50, Yolk of Egg, by volume, 6 50, Tragacanth, in powder, 0 25, Elixir of Gluside, 0 75, Simple Tincture of Benzoin, 0 75, Spirit of Chloroform, 3, Essential Oil of Bitter Almonds, 0 10, Distilled Water, *q s* to make 100

Dose—2 to 8 fl drms = 7 1 to 28 4 cc

This preparation can be medicated with any desired salt by dissolving such salt in the Water previous to making the emulsion, **Cod-Liver Oil Emulsion with Glyceriophosphates**. A Cod-Liver Oil with Hypophosphites are soluble in the Oil should be added to it before emulsification

Emulsum Olei Morrhue—Cod-Liver Oil, 500 cc, Gum Acacia, in powder, 125 grammes, Syrup, 62 5 cc, Oil of Bitter Almonds, 12 minims, Water, *q s* to make 1000 cc—*Proposed Canadian Addendum, P J '99*, n 231

This has been incorporated in the *B P C* as follows —

Emulsio Olei Morrhue—Cod-Liver Oil, 50, Gum Acacia, in fine powder, 12 50, Syrup, 6 25, Oil of Bitter Almonds, 0 10, Distilled Water, *q s* to make 100—*B P C*

Cod-Liver Oil, 50, Powdered Acacia, 12 5, Solution of Gluside, 0 75 or Syrup of Tolu, 10, Water, *q s* to make 100 Flavouring as desired—*Canadian Formulary* 1905, *C F* 1908 changed the quantity of Solution of Gluside to 0 7, but otherwise the formula is the same as 1905

Emulsum Olei Morrhue—Cod-Liver Oil, 50, Acacia in fine powder, 12 5, Syrup, 10, Oil of Gaultheria, 0 4, Water, *q s* to make 100 cc

The Oil may be replaced if desired by a suitable quantity of Oil of Bitter suitable flavouring—*U S P*

Emulsion d'Huile de Foie de Morue.—Put into a bottle 140 of Cod-Liver Oil, 4 drops of Essence of Almonds, 60 of Syrup, and 40 of Orange Flower Water, boil for 20 minutes 5 of Carrageen in sufficient quantity of water to obtain 220 of decoction with pressure through a cloth, and reduce the liquid by means of to 160, and pour the hot liquid into the bottle containing the other ingredients, agitate for 5 minutes, and then from time to time until cold, all by weight except the Essence of Almonds—*Fr Codex*

Foreign Pharmacopœias—Belg (*Tecoris Aselli Olei Emulsio*), Carrageen 10, Water 500, make 450 parts of decoction Mix, Tragacanth 1, Cod-Liver Oil 500, Anethol 2, Acetic Ether 1, Oil of Bitter Almonds 0 50 Add this mixture to the decoction and 50 parts of Glycerin Span, (*Emulsion de Aceite de Hígado de Bacalao*), Oil of Bitter Almonds 0 25, Carrageen 10, Glycerin 120, Cod-Liver Oil 500, Water to 1000

EMULSUM OLEI MORRHUE CUM HYPOPHOSPHITIBUS.—Cod-Liver Oil, 50, Acacia, in fine powder, 12 5, Calcium Hypophosphite, 0 5, Sodium Hypophosphite, 0 5, Oil of Gaultheria, 0 4, Water, *q s* to make 100—*U S P*

The Oil of Gaultheria may be replaced if desired by a suitable quantity of Oil of Bitter Almond or other suitable flavouring—*U S P*

Emulsio Olei Morrhue cum Vitæ—Cod-Liver Oil, 8 fl oz, Yolk of Egg 8 grains, Solution of Saccharine (15 pc), 1 fl drms, Simple Tincture of Benzoin, 1 fl drms, Spirit of Chloroform, 4 fl drms, Essential Oil of Almonds, 8 minims, Sodium Hypophosphite 60 grains, Calcium Hypophosphites, 60 grains, Sherry, 15 fl oz, Wine 15 fl oz, add the Hypophosphites in the Wine, place the mixture in a bottle with a little Oil, then add the yolks of Eggs, shake well, add Wine and Oil alternately until quantity is made up—*A P C*

Emulsio Olei Morrhue cum Hypophosphitibus—Cod-Liver Oil, 50, Yolk of Egg by volume, 6 50 Tragacanth, in powder, 0 25, Elixir of Gluside,

0.75, Simple Tincture of Benzoin, 0.75 Spirit of Chloroform, 3, Oil of Bitter Almonds, 0.10, Sodium Hypophosphite, 0.75, Calcium Hypophosphite, 0.75, Distilled Water, *qs* to produce 100 — *B P C*

It is sometimes made with Sherry instead of Distilled Water, and should then be distinguished as *Emulsio Olei Morrhue cum Hypophosphitibus et Vino*

Foreign Pharmacopœias — *Span* (*Emulsion de Acete de Hgado de Bacalao con Hipofosfitos*), Oil of bitter Almonds 0.25, Calcium Hypophosphites 5, Soda Hypophosphites 5, Carayeen 10, Glycerin 120, Cod Liver Oil 500, Water to 1000 *Swiss* (*Emulsio Olei Jecoris*), Cod Liver Oil 1000, Gum Arabic 10, Tragacanth 10, White Gelatin 2, Calcium Hypophosphites 5, Soda Hypophosphites 5, Sugar, 0.2 Cinnamon Oil 4 drops, Alcohol 50, Orange Water 40, Water 878 *US* (*Emulsio Olei Morrhue et Emulsio Olei Morrhue cum Hypophosphites*), *see above* *Mex* (*Emulsion de Acete de Bacalao*), Tragacanth 4, Glycerin 50, Cod Liver Oil 250, Oil of Bitter Almonds 0.25, Calcium Hypophosphites 5, Sodium Hypophosphites 5, Water 200

HUILE DE FOIE DE MORUE CRÉOSOTÉE — *Cicosoto*, 1, Cod Liver Oil, 99 — *Fr Codex*

HUILE DE FOIE DE MORUE PHOSPHORÉE — Cod Liver Oil, 497.5, Phosphorated Oil (1 p c), 2.5 — *Fr Codex*

MORRHUOL — Cod Liver Oil treated first with aqueous solution of Sodium Bicarbonate to remove the acids, then agitated with Alcohol (90 p c), which on evaporation yields Morrhual. Brown Oil yields $4\frac{1}{2}$ to 6 p c, the straw coloured $2\frac{1}{2}$ to 3 p c — *Y B P '86*, 231, *P J '97*, 11438

Proposed as a substitute for Cod Liver Oil, but without the Carbo hydrates, and, owing to its small bulk, is adapted for administration in capsules

Dose — 3 grains = 0.2 gramme

MOSCHUS.

MUSK

FR, *MUSC*, *GER*, *BISAM*, *ITAL*, *MUSCHIO*, *SPAN*, *ALMIZCLE*

The dried secretion from the preputial Follicles of *Moschus moschiferus*, L. Dark reddish-brown grains or masses of grams, somewhat unctuous to the touch, and possessing a peculiar penetrating persistent odour and a somewhat bitterish taste

Dan, should be practically free from moisture, and yield not more than 8 p c of ash, *US*, 8 p c of ash

Commercial samples contain large quantities of moisture (about 30 p c)

The Musk deer is a native of the mountainous regions of Central Asia. Musk is imported from China and India

Medicinal Properties — A diffusible stimulant and antispasmodic. Used in hysteria and spasmodic asthma, and as a stimulant in the prostration of diseases such as typhoid

Dose — 5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes — Usually prescribed in a mixture or in pills. *See formulas given below*

Not Official — *Mistura Moschi*, *Moschus Fasicatus*, *Pilula Moschi*, and *Tinctura Moschi*.

Foreign Pharmacopœias — Official in all except *Austr*, *Belg*, *Dan*, *Ger* and *Ital*, *Fr* (*Musc*), *Port* (*Almizcar*), *Mex* and *Span* (*Almizcle*)

Descriptive Notes—Musk enters commerce in two forms, viz, either in the original sacs or "pods," or in a granular powder known as Giam Musk, consisting of the dried contents of the sacs. The best kind is that known as Tonquin or China Musk, it occurs in oval, plano-convex sacs, covered on the convex side with long hairs, and has a small nearly central orifice around which bristly hairs converge. A pod contains on the average about 40 to 45 grammes of Giam Musk. It is imported *via* Canton. Yunnan Musk comes next in value, it comes *via* Shanghai, the sacs are more rounded and less oval, with crowded short hairs, and the orifice is usually stopped with a plug of rice straw. The Assam or Nepaul Musk occurs in sacs that have a rounder outline and a larger orifice, or are more spherical ('bally'), with the inner surface partly filled with a grisly substance and contains proportionately less grain than the other kinds. Cabardine, Russian or Siberian Musk, so called from the name Kabarga given to the animal in the Altai Mountains, is of a narrower oval form than the Tonquin Musk, more flattened, and the hairs are drier and the flat or ventral surface often has a white efflorescence. Its odour is weaker and slightly ammoniacal, and it is the least valuable kind. Musk is sometimes falsely packed, when stitches can be seen round the margin, especially if placed in lukewarm Water, or adulterated by the insertion of particles of lead, etc. If the hairs do not cling to the orifice the sac is of artificial character. The sacs are usually tested by probing with a long pin and the odour judged by the Musk attached to the pin when withdrawn, but it is necessary to use a fresh pin for each pod or the odour of a good pod or sac may be communicated to an adulterated one. The grains are officially stated to be 'contained in an oval sac $1\frac{1}{2}$ to 2 in (37 to 50 mm) in diameter, nearly smooth on one side and covered on the outer side with hairs concentrically arranged around a nearly central orifice,' this description applying to Tonquin or China Musk pods. Artificial Musk that can be used as a perfume has recently been the subject of several patents.

Tests.—Musk is officially required to be free from impurities of an earthy nature, and should not leave more than 8 p.c. of ash when ignited with free access of air. Musk contains a considerable proportion of moisture, amounting to about 30 p.c., the *USP* requires that the moisture should not exceed 15 p.c. and that the ash should not amount to more than 8 p.c. The moisture is determined by drying the Musk over Sulphuric Acid.

Not Official.

MISTURA MOSCHI—Musk, 3, Gum Acacia, 3, Sugar, 3, Rose Water, 160, triturate the Musk with the Sugar, then with the Gum Acacia, add the Rose Water gradually.

Dose—1 to 2 fl. oz. = 28.4 to 56 S.c.c.

An emulsion (1 in 100) is **Official** in Swed.

MOSCHUS EXSICCATUS—Musk which has been dried over Strong Sulphuric Acid. It keeps better than that which is usually supplied as 'giam Musk.' It is easily made into pills with Dispensing Syrup or 'Diluted Glucose.'

PILULA MOSCHI—Musk, 12, Powdered Gum Acacia, 3, Powdered Liquorice, 3

Dried Musk, 12, Powdered Gum Acacia, 1, 'Diluted Glucose,' *qs*

TINCTURA MOSCHI—Musk, 5, Alcohol (95 p c), 45, Water, 15 Alcohol (45 p c), *qs* to produce 100 Triturate the Musk with the Water a little at a time until a smooth mixture is obtained, transfer to a bottle, and allow it to stand for 24 hours, add the Alcohol and macerate the mixture for 6 days, occasionally shaking it, then filter through a plain paper filter and, when the liquid has drained off completely, pass enough Alcohol (15 p c) through the filter to make 100 of Tincture—*U S P*

This has been incorporated in the *B P C* as follows—

Musk, 5, Alcohol (90 p c), 50, Distilled Water, 15 Alcohol (15 p c), *qs* to produce 100—*B P C*

Fr, Mex and Port—Musk, 1, Spirit, 10

Dutch, Russ and Swiss—Musk, 1, Spirit, 25, Water, 25

Span—Musk, 1, Spirit, 25

U S—Musk, 5, Water, 45, Alcohol, 45, Diluted Alcohol to measure 100

All by weight except U S

Not Official

MYROBALANUM

The dried immature Fruits of *Terminalia Chebula*, Læv., commonly known as *Chebulic myrobalans*. A local astringent

Dose—30 to 60 grains = 2 to 4 grammes

UNGUENTUM MYROBALANI—Myrobalans, in very fine powder, 1, Benzoated Lard,* 4—*Ind* and *Col* Add

UNGUENTUM MYROBALANI CUM OPIO Myrobalan Ointment, 92½, Opium, in powder, 7½—*Ind* and *Col* Add

MYRISTICA.

NUTMEG

FR, MUSCADE DES MOUQUES, GER, MUSKATNUSS, ITAL, NOCI MOSCATA, SPAN, NUEZ MOSCADA

The dried Seed of *Myristica fragrans*, Houtt, divested of its testa.

It is cultivated in the Banda Islands of the Malayan Archipelago and imported from Sumatra and the Molucca Islands, and occasionally from the West Indies and the Seychelles

Medicinal Properties—Aromatic, stimulant, and carminative. Frequently used to cover the taste of Rhubarb and other medicines. The expressed and volatile Oils have been much used in chronic rheumatic pains and in lotions for the hair. In large and poisonous doses it resembles the action of *Cannabis Indica*.

Severe toxic symptoms followed on taking a whole nutmeg, grated, in a wineglassful of Gin, to procure abortion—*L* '02, 1 1035, 1798

Paper on nutmeg poisoning by Cushman, he notes the scanty recognition of this in English medical literature, and narrates the death of a boy after eating two nutmegs—*B M J* '08, 1 387 and *L* '08, 1 495

Dose—5 to 15 grains = 0.32 to 0.1 gramme

* *Adeps Induratus* (Lard deprived of a portion of its Oil by pressure) may be employed in India and the Colonies when prevailing high temperatures render it necessary. See *Appendix I, Ind*, and *Col, Add*.

Prescribing Notes—The Oil may be given on Sugar, or in pill with Liquorice powder and Soap, see p 897

Official Preparation—Oleum Myristicæ Used in the preparation of Pulvis Cretæ Aromaticus, Spiritus Armoiacæ Compositus and Tinctura Lavandulæ Composita, of the Oil, Spiritus Myristicæ Used in the Spiritus Ammoniac Aromaticus, Tinctura Guaiacæ Ammoniata, Ammoniata and Pilula Aloes Socotinae Of the Spirit, contained in Mistura Ferri Composita

Not Official—Oleum Myristicæ Expressum (Myristicæ Adops)

Foreign Pharmacopœias—Official in Austr, Dutch, Ger, Russ, Swed and Swiss (Semen Myristicæ), Belg and Hung (Nux Moschata), Fr (Muscadées Moluques), Ital (Noce Moschata), Port (Noz Moschada), Mex and Span (Nuez Moscada), US (Myristica) Not in Dan Jap or Norw

Descriptive Notes.—Nutmegs, freed from the endocarp or shell, are imported into this country from Penang, Singapore, Bombay, and the West Indies The Dutch formerly treated the Nutmegs with Milk of Lime with the view of preventing germination, and so securing a monopoly, but avowedly to protect them from the attack of insects, but exposure to the sun for a week effectually prevents germination. The Chinese prefer to import them in the shell, dried till they rattle inside, in spite of the increase of cost of freight, the aroma of the seed is best preserved in this manner Nutmegs are sorted according to size, and may vary from 66 to 132 to the lb, those from 66 to 84 being of good size and fetching, if in good condition, the highest price The smaller and defective or irregularly formed Nutmegs are used for distilling the Oil, and for expressing the fixed Oil known as Nutmeg Butter, and erroneously as Oil of Mace, Nutmeg Butter is also imported from Sumatra Nutmegs are obtusely oval or rounded, and vary very much in size, those of good quality are 1 to $1\frac{1}{2}$ in long (25 to 31 mm) and $\frac{7}{8}$ in (22 mm) broad (rarely exceed 1 in (25 mm) in length, BP) The surface is furrowed and veined, with a circular scar at the broad end, the transverse section shows ruminated Albumen caused by the infolding layer of the light brown perisperm, it is easily cut, and has a waxy lustre, a characteristic odour, and a slightly bitter, aromatic taste The microscopical characters of Nutmegs are the dark brown tabular cells of the perisperm and the polyhedral cells of the endosperm, some containing starch grains with a central hilum, often in groups of 4 to 8, others a large monohedral crystalline and others filled with brown oleoresinous coloring matter Malabar Nutmegs (*Myristica Malabarica*, Lam) which are sometimes offered for sale have practically no aroma Macassar Nutmegs (*Myristica argentea*, Warb) imported from New Guinea via Macassar have a faint Nutmeg flavour, but are much more acrid Both of these are longer than true Nutmegs, exceeding an inch and a half as a rule

Mace, the arillus of Nutmegs, is sold separately When fresh it is of a bright crimson red colour, but fades to bright orange brown The distinctive microscopical characters of true Mace are the polymorphous grains of amylopectin in the parenchymatous cells, which are coloured brownish red by solution of Iodine in Iodide of Potassium, and the large

elongated thick walled quadrangulum, epidermal cells, often with pointed ends Powdered Mace is frequently adulterated with Bombay Mace, the latter can be detected by the rounded cells containing a dark yellow colouring matter, often free, and by the red flocculent precipitate given in an alcoholic extract of the Mace by the solution of Lead Acetate, if the Mace is genuine, only a milky white turbidity is produced Turmeric, which is not likely to be used, gives a similar precipitate, but filtering paper dipped in the alcoholic solution and dried gives with Boric Acid an orange red or red brown colour, which Bombay Mace does not Bombay Mace also contains a fat which has different chemical characters from that of the true Mace *Myristica fatua*, Houtt, and *M. cinnamomua*, King, appear to be the only species resembling in flavour the true Nutmeg

Tests—Nutmeg when incinerated with free access of air yields from 2 to 3 p.c. of ash, and the figure should not exceed 4 p.c. It yields from 9 to 15 p.c. of volatile Oil, which should answer the tests given under *Oleum Myristicæ*, it also contains from 30 to 40 p.c. of fixed Oil

Preparations

OLEUM MYRISTICÆ OIL OF NUTMEG

A colourless, or pale yellow, mobile liquid, possessing a characteristic odour and a spicy taste, distilled from Nutmeg

It darkens in colour and becomes viscid by oxidation on exposure to air It should therefore be kept in well closed glass bottles of a dark amber tint and protected as far as possible from exposure to the air and light

Solubility—In all proportions of Absolute Alcohol, 1 in 4½ of Alcohol (90 p.c.), sparingly in Alcohol (60 p.c.)

Dose—½ to 3 minims = 0.03 to 0.18 cc

Foreign Pharmacopœias—Official in Austr, Dutch, Ger, Hung, Russ and Swiss (*Oleum Macidis*), Belg (*Essentia Myristica*), Jap (*Oleum Myristicæ Ætherum*), Norw and Swed, (*Ætherioleum Macidis*), Port (*Essencia de Noz Moschada*), U.S. (*Oleum Myristicæ*)

Tests—The Oil of Nutmeg has a sp. gr. of 0.865 to 0.930 The official figures are 0.870 to 0.910, the *USP* 0.884 to 0.924 at 25° C (77° F), the *Ph* 0.890 to 0.930 It is officially required to form a clear solution with its own volume of a mixture of equal parts of Absolute Alcohol and Alcohol (90 p.c.), the *USP* states that it is soluble in an equal volume of Alcohol (94.9 p.c.) and also soluble in 3 volumes of Alcohol (90 p.c.), the *Ph* states that it dissolves in 3 parts of Alcohol (90 p.c.) The Oil is optically dextrogyrate, the rotation being from +7° 52' to +28 in a 100 mm tube The *BP* requires that the Oil when evaporated on a water-bath shall not leave a residue which crystallises on cooling, which indicates the absence of concrete Oil of Nutmeg The *USP* states that when 2 or 3 cc of Oil are evaporated on a water bath no residue which crystallises on cooling should be left The Pharmacopœia test for concrete Oil of Nutmeg has been somewhat adversely criticised,

attention is drawn (*P J* '01, 1 328) to the necessity for modification of the pharmacopœial description. The requirement that the Oil should yield, when evaporated on a water-bath, no residue which crystallises on cooling, necessitates the attachment of some qualifying phrase, such as 'by fractionation,' and the Pharmacopœia description as the 'Oil distilled from Nutmeg,' otherwise the tests given in the *B P* would seem to require modification.

The product is a mixture of fatty Oil which is carried over mechanically, and is supposed to cause an unpleasant preponderance of the Nutmeg over the lemon flavour in *Sal Volatile*.

The Oil contains Dextro- and Lævo-Pinene, Dipentene, Myristicol, Myristicin, Myristinic Acid, and a Phenol-like substance which yields an emerald green with Ferric Chloride T S.

SPIRITUS MYRISTICÆ SPIRIT OF NUTMEG

Oil of Nutmeg, 1, Alcohol (90 p c), *qs* to produce 10

Should be clarified if necessary, by means of Talc

B P 1885 was 1 in 50

Dose —5 to 20 minims = 0.3 to 1.2 c c

Tests—Spirit of Nutmeg has a sp gr of about 0.833, it contains about 0.5 p c w/v of total solids, and about 86.0 p c w/v of Absolute Alcohol.

Not Official

OLEUM MYRISTICÆ EXPRESSUM *Syn.* MYRISTICÆ ADIPS—A concrete Oil, of a firm consistence and orange colour, obtained from Nutmeg by expression and heat.

Foreign Pharmacopœias—Official in Aust. (*Ol Myristicæ Expressum*), Dutch and Swiss (*Oleum Myristicæ*), Fr (*Beurre de Muscade*), Ger (*Ol Nucistæ*), Mex (*Manteca 'O Aceite concreto de Nuez Moscada*), Port (*Oleo de Noz Moscada*). Not in the others.

MYRRHA.

MYRRH

Fr, MYRRHE, GER, MYRRHE, ITAL, MIRRA, SPAN, MIRRA

Small, irregular, brownish-yellow, or reddish-brown, rounded fragments, or tears, or masses of them, having a dusty appearance on the surface, and possessing a strong characteristic aromatic odour. It is a Gum Resin obtained from the Stem of *Balsamodendron Myrrha*, and probably other species.

Collected in Somaliland and South-eastern Arabia.

Myrrh contains from 57 to 59 p c of Gum, a neutral Resin, a soft Resin and two acid Resins, and from 7 to 8 p c of an ethereal Oil.

Solubility—Myrrh contains from 40 to 65 p c of Gum soluble in Water, the remainder, consisting of Resin, is mostly soluble in Alcohol.

Medicinal Properties—Stomachic and carminative, expectorant. Locally to aphthæ of mouth and spongy gums.

Prescribing Notes—*The Tincture mixed with Water (1 to 24) is used as a gargle, but the addition of Mucilage of Gum Acacia is often necessary, also mixed with Solution of Borax as a mouth wash*

Official Preparation—*Tinctura Myrrhae* Contained in *Decoctum Aloes Compositum*, *Mistura Ferri Composita*, *Pilula Aloes et Myrrhae*, *Pilula Galbani Composita*, and *Pilula Rhei Composita*

Not Official—*Gargalisma Myrrhae (Squue)*

Foreign Pharmacopœias—Official in all except Hung

Descriptive Notes—Myrrh as imported is usually mixed with the Gum Resins of allied species of *Balsamodendron*, which need to be removed. None of these have the strong bitterness of Myrrh, but most of them are acrid. Bissabol, the product of *Balsamodendron Erythæum*, var. *glabrescens*, Engl., has a distinct odour and taste, but much resembles Myrrh in colour and in the presence of white streaks of gum. It does not give the violet colour with the official test for Myrrh (*P J* (4) xiii 666–7). It is known in commerce as ‘Opopanax,’ and is the source of the ‘Oil of Opopanax’ used in perfume. True Opopanax Oil has a strong flavour of Celery or Lovage, and is the product of an umbelliferous Gum Resin imported from Persia. Indian and African Bdelliums are acrid, but have neither the taste nor odour of Myrrh, and have not white streaks of gum, the former has a taste like cedar, they have a dull fracture and are tougher and less easily broken than Myrrh.

Pure Myrrh is characterised by its distinctive odour and flavour and its bitter taste, and by assuming a violet colour when moistened with Nitric Acid, which the false Myrrhs do not present when so treated. It varies much in size and form, but is usually reddish-brown externally, with a powdery surface, and when broken either presents a dull uniform resinous fracture, often somewhat translucent, or in other pieces there occur distinct white streaks, indicating gum. Crude or unpicked Myrrh should not be used in pharmacy, as it contains so much foreign matter that its price is only one-third of that of selected Myrrh. Pieces in which coiled drops of thickened oil have exuded on the surface will be richer in aroma and contain more oil. Pieces containing white streaks are more suitable for emulsion, and those with little gum are preferable for tincture, the gum being insoluble in Alcohol. The gum left after making the tincture makes a good adhesive mucilage when dissolved in water.

Tests—Myrrh is officially required to yield a violet colour when moistened with Nitric Acid. This test has been modified (*P J* '01, ii 666) to 0.5 of a gramme of coarsely powdered Myrrh mixed with 10 c.c. of Ether and shaken occasionally for 10 minutes yields a filtrate, 2 c.c. of which should yield on evaporation a residue which is coloured violet with the vapour of Nitric Acid. No percentage of matter soluble or insoluble in Alcohol (90 p.c.) is given nor is any reference made to the percentage of ash. The *U S P* states that in Alcohol (94.9 p.c.) it yields a brownish-yellow tincture, acquiring a purplish-red tint on the addition of Nitric Acid, but no percentage of matter soluble or insoluble in Alcohol is recorded. The German Pharmacopœia requires that when 1 gramme of powdered Myrrh is

shaken with 2 or 3 grammes of Ether, the liquid filtered and the yellow filtrate treated with Bromine vapour, a reddish-violet coloration is produced. This Bromine reaction is usually considered to be unreliable. A test which is stated to distinguish between Heerabol and Bissabol Myrrh is as follows—Mix 6 drops of a 1 in 15 Petroleum Ether extract of the Myrrh with 3 cc of Glacial Acetic Acid, and float the mixture carefully on the surface of 3 cc of Saturated A. C., a rose-red coloration forms at the point of contact of the two liquids.

100 parts of Myrrh when completely exhausted with boiling Alcohol (90 p.c.) are required by the *P.G.* to leave a residue which upon drying shall not yield more than 70 parts, the percentage of ash shall not amount to more than 6 p.c. Myrrh is stated (*C.D.* '00, 1 101) to be easily obtainable of good quality, it should leave when exhausted by Alcohol (90 p.c.) not more than 60 p.c. of insoluble residue, and the ash should not exceed 5 p.c. and should be almost entirely soluble in dilute Hydrochloric Acid.

Dieterich suggests a limit of not more than 70 p.c. of matter insoluble in Alcohol and that the percentage of ash should not be more than 10 p.c., and also suggests the inclusion of figures for the Acid, Ester and Saponification values of the Gum Resin, and gives for Heerabol Myrrh containing 20 p.c. of matter soluble in Alcohol an Acid value of 25.48, an Ester value of 204.12, and a Saponification value of 229.60.

A sample of Gum Myrrh Elect examined in the author's laboratory gave 49.94 p.c. of matter insoluble in Alcohol (90 p.c.), 3.75 p.c. of ash. It contained 50.06 p.c. of matter soluble in Alcohol (90 p.c.) and possessed an Acid value of 19.6, an Ester value of 118.44 p.c. and a Saponification value of 138.04 p.c. Other samples of the Gum Resin examined for percentage of ash only gave from 3.3 to 4.6 p.c. Samples of powdered Myrrh gave from 4.7 to 6.2 p.c. of ash. Two samples of the powder gave 45.92 and 45.69 for the Acid value 86.24 and 91.51 for the Ester value, and 132.16 and 137.20 for the Saponification value. They contained respectively 6.2 p.c. and 6.05 p.c. of ash. They were unfortunately not examined for the percentage of matter soluble in Alcohol (90 p.c.). A sample of powder yielded 4.5 p.c. of ash and yielded 56.90 p.c. of matter soluble in Alcohol (90 p.c.).

Preparation

TINCTURA MYRRHÆ. TINCTURE OF MYRRH

Myrrh, 1, Alcohol (90 p.c.), *q.s.* to yield 5 (1 in 5)

B.P. 1885 was 1 in 8

Dose— $\frac{1}{2}$ to 1 fl. dr. = 1.8 to 3.6 cc

Foreign Pharmacopœias—Official in Aust., Belg., Dan., Dutch, Ger., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S., 1 in 5. All by weight except U.S. Not in Fr. or Hung.

Tests—Tincture of Myrrh has a sp. gr. of 0.845 to 0.855, it contains from 4 to 6 p.c. w/v of total solids, averaging about 5 p.c. w/v, but this figure must necessarily depend upon the amount of

matter soluble in Alcohol (90 p c) contained in the Gum Resin used, it also contains about 85 p c w/v of Absolute Alcohol. A few drops of the Tincture evaporated on a water bath yield a residue giving a violet coloration when moistened with Nitric Acid.

Not Official

GARGARISMA MYRRHÆ (*Squire*)—Tincture of Myrrh, 1, Honey, 1, Infusion of Roses, 18.

This has appeared in *Squire's Companion* since 1861 and is now incorporated in the *B P C*.

TINCTURE OF MYRRH AND BORAX—*See BORAX*.

Not Official

NAPHTHALINUM

NAPHTHALENE

$C_{10}H_8$, eq 127.10

Crude Naphthalene is a hydrocarbon, crystallising from coal tar. When purified by sublimation it occurs in white micaceous scales, with a characteristic odour, melting at $80^{\circ}C$ ($176^{\circ}F$).

Solubility—Insoluble in Water, soluble 1 in 25 of Alcohol (90 p c), 1 in $1\frac{1}{2}$ of Chloroform, 1 in 3 of Ether, 1 in $7\frac{1}{2}$ of Oil of Turpentine, 1 in 5 of Olive Oil, slightly soluble in Glycerin.

Medicinal Properties—Antiseptic, the fine powder is dusted over ulcers and wounds, and is useful for disinfecting cavities. It is given as an intestinal disinfectant. A parasiticide in scabies, as 10 p c solution in Olive Oil, or as an Ointment.

Crude Naphthalene in balls and other shapes is used to protect furs and woollen articles from moths.

Dose—Usual dose, 2 to 5 grains = 0.13 to 0.32 gramme, every four or six hours. Larger doses have been given, but are apt to upset digestion, and in some cases to produce toxic symptoms.

Prescribing Notes—*It has a nauseous taste and odour, when given internally it may be enclosed in a cachet or capsule, or made into pills with 1 of Compound Tragacanth Powder to 12, and massed with Diluted Glucose.*

Foreign Pharmacopœias—Official in Aust, Dutch, Ger, Ital, Jap, Mex, Russ, Swed, Swiss and U.S. Not in the others.

Tests—Naphthalene melts at $80^{\circ}C$ ($176^{\circ}F$) and boils at $218^{\circ}C$ ($421^{\circ}F$), it volatilises slowly at the ordinary temperature and completely with further heat, it burns with a luminous smoky flame. It should not possess an acid reaction to blue Litmus paper moistened with Water, indicating the absence of free acids, e.g., Sulphuric Acid. It should dissolve colourless in warm concentrated Sulphuric Acid if quite pure, but a decided pinkish tint is observed if the sample contains 1 p c of impurity, the coloration becoming a deeper pink or even brown the larger the proportion of foreign matter present. 0.5 of a gramme when ignited with free access of air should leave no weighable residue, indicating the absence of mineral impurity.

NAPHTHALINUM PRÆCIPITATUM—A fine powder, obtained by dissolving the crystals in hot Alcohol, and pouring into a quantity of cold Water, Recommended as less irritating than the powdered crystals.

PULVIS NAPHTHALINI (*Rosbach*)—Purified Naphthalene, 75 grains Sugar, 75 grains, Oil of Bergamot, $\frac{1}{2}$ minim, divide into 20 powders. In vesical catarrh.—*L.* '85, 1, 360.

NAFTALAN—A dark, greenish-black, unctuous substance, insoluble in Water, soluble in Ether and in Chloroform, an extraction product of a naphtha from the Caucasus containing about 96 p.c. Soap. Used in various skin diseases, and stated to be a good vehicle for the application of antiseptic preparations—*L* '99, 1 1234, *B M J E* '99, 1 92, *P J* '01, 1 124, *B M J E* '05, 1 40. It may be applied (*B M J E* '05, 1 64) as an ointment or in the form of suppositories.

Naphthalini Tetrachloridum—Glistening, white crystals, insoluble in Water.

Dose—2 to 10 grains = 0.13 to 0.65 gramme

NAPHTHOL.

BETA-NAPHTHOL

BETA-MONO-HYDROXY-NAPHTHALENE

$C_{10}H_8O$, eq. 142.98

Fr., NAPHTHOL, GER., BETANAPHTHOL, ITAL., NAFTALOLO

White, or pale yellowish, lustrous, crystalline laminae or as a white or almost white crystalline powder, having a faint odour suggestive of Phenol, and a sharp, biting but not persistent taste.

It is described in the *USP* as a monatomic Alcohol, occurring in coal-tar, but usually prepared from Naphthalene, the *BP* states that it is derived from Naphthalene-sulphonic Acid.

It should be kept in well-closed bottles of a dark amber tint and protected as far as possible from the air.

There are two isomeric Naphthols, Alpha-Naphthol and Beta-Naphthol, bearing the same relation to Naphthalene as Phenol does to Benzol.

Solubility.—Nearly insoluble in Water, soluble 1 in 2 of Alcohol (90 p.c.), 3 in 4 of Ether, 1 in 24 of Chloroform, 1 in 12 of Olive Oil, 1 in 40 of Glycerin.

Aqueous solution of Boric Acid will dissolve comparatively small quantities of Naphthol.

Medicinal Properties.—Disinfectant, intestinal antiseptic. Given in summer diarrhoea of children, and in typhoid and intestinal dyspepsia, prolonged administration, especially of large doses, may lead to nephritis. Used in parasitic skin diseases and in chronic eczema in form of ointment.

Dose—3 to 10 grains = 0.2 to 0.65 gramme

Prescribing Notes.—Given in cachets or pills. A good pill can be made by adding a small quantity of Compound Powder of Tragacanth and Dispensing Syrup, or 'Diluted Glucose' q.s. Also administered dissolved in Oil, which is then emulsified. It can be made into an Ointment with Lard, Soft Paraffin or Lanolin Ointment, for Kaposi's Ointments, see below.

When no prefix is attached to the name, Beta-Naphthol should be used. The name is also written *Naphthol*.

Should be kept in dark amber-tinted well stoppered bottles.

Not Official.—Lassar's Itch Remedy, Naphthol-Camphor, Pommade Naphtholée, Parogenum Naphtholæ, Unguentum Naphtholi, Unguentum Naphtholi Compositum, Vaselineum Naphtholi, Asaprol, Benzonaphthol, Betol, Epicarin, Quinaphthol.

Foreign Pharmacopœias —Official in Austr, Belg, Dan, Dutch, Fl, Ger, Hung, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U S

Tests —Beta Naphthol when pure melts at 122° C (251 6° F) and boils at 286° C (546 8° F), the *B P* gives the above m p but not the boiling point, the *U S P* and *P G* give the above-mentioned m p and boiling point

A hot saturated aqueous solution yields a blue fluorescence with one drop of Ammonia Solution, if some Chlorine Solution be added to a cold saturated aqueous solution a white turbidity is produced, and on the addition of Ammonia Solution a green or brownish coloration is produced. In performing this test the *U S P* uses in the place of Chlorine Water a solution of Chlorinated Lime and requires that it should produce a yellow colour. The *B P* requires that 10 c c of a 1 p c boiling aqueous solution should yield a white precipitate becoming brown upon the addition of 10 drops of 3 p c aqueous Ferric Chloride Solution. A saturated aqueous solution yields no colour upon the addition of a few drops of Iodine T S followed by an excess of Sodium Hydroxide Solution. It is distinguished from its isomer Alpha-Naphthol by the m p, Alpha-Naphthol melting at 95° C (203° F), by the reaction with Chlorinated Lime Solution, Alpha-Naphthol giving a dark violet colour, by the test with Ferric Chloride Solution, Alpha-Naphthol giving a violet coloration with the latter reagent, and by the test with Iodine Solution and excess of Sodium Hydroxide Solution, which produces an intense violet coloration with Alpha-Naphthol.

The more generally occurring impurities are free acid, Naphthalene, organic impurities, and impurities of a mineral or inorganic nature. It should possess a neutral reaction towards Litmus paper previously moistened with Alcohol (90 p c). Naphthalene and organic impurities may be detected by the Ammonia test given in small type below. 0.5 of a gramme when ignited with free access of air should leave no weighable residue.

Residue —When heated it readily sublimes, and is volatilised from its aqueous or alcoholic solution with the vapour of Water or Alcohol, *U S P*. It should leave no residue on ignition, *B P* and *U S P*. 0.2 gramme should not leave a weighable residue after ignition, *P G*.

Ammonia —Beta Naphthol should be soluble in 50 parts of Ammonia Water without residue, and the solution should not have a deeper colour than pale yellow, *U S P* and *P G*.

Chlorinated Lime —A cold saturated aqueous solution should not show a violet colour with solution of Chlorinated Lime, *P G* and *U S P*, the latter stating that the colour should be pale yellow on the addition of Chlorinated Lime to an aqueous solution.

Not Official

UNGUENTUM NAPHTHOLI (Kaposi's Ointment) —Beta Naphthol, 60 grains, Prepared Lard, 1 oz.
Beta-Naphthol, 10, Lard, 90 —*B P C*

POMMADE NAPHTHOLÉE —Beta-Naphthol, 10, Vaseline, 90 —*F*

UNGUENTUM NAPHTHOLI COMPOSITUM (Kaposi) —Naphthol, 15, Prepared Chalk, 10, Soft Soap, 50, Lard, 100.

Official in Austr, β -Naphthol, 10, Precipitated Chalk, 5, Soft Soap, 28; Lard, 37

LASSAR'S ITCH REMEDY—Beta-Naphthol, 0.25, Peru Balsam, 10, Spirit Soap, 25—*Hager*

VASOLIMENTUM NAPHTHOLI—Beta-Naphthol, 10, Vasoliment Liquid, 90—*Hager*

Parogenum Naphtholis *Syn* Naphthol Vasoliment—Naphthol, 10, Parogen, *qs* to produce 100—*B P C*

ASAPROL ABRASTOL (Calcium B:ate) —A white powder, soluble in Water. *H* pyretic and analgesic, in sciatica, muscular and chronic rheumatism and in chronic nephritis.—*T G* '93, 182, '94, 252, *Pr* 111 52, *MA* '95, 8, *Y B T* '94, 462, '95, 159

Dose—5 to 15 grains = 0.32 to 1 gramme

BENZONAPHTHOL $C_{10}H_7O_2$, eq 223.7. Prepared by the action of Benzoyl Chloride on Beta-Naphthol, tasteless powder, almost insoluble in Water and Ether, soluble in Chloroform. Intestinal antiseptic, and disinfectant. Has been found useful in typhoid—*Pr*, 11 213. In tropical dysentery—*L* '95, 11 169, *P J* '95, 11 238

Dose—5 to 15 grains = 0.32 to 1 gramme

Official in Fr, Mex, Spain and Swiss

Tests—Benzonaphthol melts at 107° to 108° C (224.6° to 226.4° F), the *Fr Codex* (1908) gives 110° C (230° F), it dissolves in Sulphuric Acid with the production of a pale yellow colour, if the Sulphuric Acid solution be diluted with Water and rendered alkaline with an excess of Ammonia Solution, green fluorescence is produced. When warmed with Potassium Hydroxide Solution it is decomposed, if the solution be exactly neutralised with dilute Sulphuric Acid, it yields with Ferric Chloride T.S. a buff precipitate. The Benzoic Acid separated from the salt should possess the characteristic tests given under Acidum Benzoicum. A small amount of Potassium Hydroxide Solution, cooled, rendered turbid by the addition of diluted Sulphuric Acid yields on the addition of Chlorinated Lime Solution a yellow colour but no dark violet coloration.

BETOL NAPHTHALOL β -Naphthol Salicylic Ester. Beta-Naphthol Salicylate β -Naphthol Salicylic Acid Ester $C_{16}H_{11}O_3$, eq 262.11. In tasteless small white crystals, or as a white, odourless, tasteless and crystalline powder insoluble in Water, soluble in Alcohol and in fixed Oils. Recommended in rheumatism, catarrhs and intestinal fermentation.

It may be prepared by the action of Phosphorus Oxychloride on a mixture of Sodium Beta-Naphthol and Sodium Salicylate.

Dose—2 to 8 grains = 0.13 to 0.52 gramme as a powder, or in pills with Glucose.

Official in Fr (Salicylate de Naphthyle- β) and *Ital*

In pencils for gonorrhoea containing 20 pc of Betol with Oil of Theobroma.

Tests—Beta-Naphthol Salicylate melts at 95° C (203° F), which figure is also given in *Fr Codex* (1908), when shaken with Water and filtered, the filtrate should be neutral in reaction towards Litmus paper. When treated with Sulphuric Acid it yields a yellow coloration and in the course of a few minutes a brown-yellow coloured solution, which on the addition of a drop of Nitric Acid changes to a brownish-green. A 1 pc solution of the salt in Alcohol (90 pc) yields with 1 drop of a strongly diluted Ferric Chloride Solution a violet coloration, but the highly diluted Ferric Chloride Solution is only rendered turbid and no coloration is produced, when 10 to 20 drops of a 1 pc solution of the salt in Alcohol (90 pc) is added to it. When heated with Potassium Hydroxide Solution it is decomposed, forming Potassium Beta-Naphthol and Potassium Salicylate. If the solution be carefully neutralised with Hydrochloric Acid it yields on the addition of Ferric Chloride T.S. a violet

coloration, the salt is also decomposed by concentrated acids yielding Salicylic Acid and Beta-Naphthol, the separated Salicylic Acid should possess the m p and answer the tests given under Acidum Salicylicum. When heated with Potassium Hydroxide Solution, cooled and slightly acidified with diluted Sulphuric Acid it yields on the addition of Chlorinated Lime a yellow but not a dark violet coloration. 0.5 of a gramme of the salt when ignited with free access of air should leave no weighable residue.

Alphol is the Salicylate of α Naphthol Ester.

EPICARIN (Beta oxynaphthyl orthoxy meta toluylc Acid) — Colourless needle shaped crystals, or a pale yellow powder, insoluble in Water, soluble in Alcohol and Ether. Employed in the form of a 10 to 20 p.c. Ointment in psoriasis, eczema and other skin affections, and in the form of a 5 to 10 p.c. alcoholic solution for seborrhoea capitis and lichen planus. — *J. L. P.* '02, 177.

The Sodium salt of the above is also known commercially.

Tests — Epicarin melts at 199°C (390.2°F), it dissolves readily in Alcohol (90 p.c.), yielding a solution which gives, on the addition of Ferric Chloride T.S. a deep blue colour, it yields when treated with concentrated Sulphuric Acid a reddish brown solution possessing a strong green fluorescence, when mixed with Potassium Hydroxide Solution and shaken with Chloroform a yellowish turbidity is produced subsequently changing to a yellowish green. When heated with free access of air it should leave no weighable residue.

NAPHTHOL-CAMPHOR — Mix 2 of Camphor with 1 of Beta Naphthol to a viscous consistency, insoluble in Water, soluble in Oils, it is strongly antiseptic. — *Hager*.

A difference of opinion appears to exist as to the harmlessness of this preparation for the treatment of certain localised tuberculous lesions. On the one hand, no grave sequels were shown to have followed 10,000 injections, and on the other, 12 deaths are reported after its use. — *L.* '04, ii 1493, *P.* '05, i 177.

QUINAPHTHOL (Quinine Beta naphthol sulphate) — A yellow crystalline powder, sparingly soluble in Water and in Alcohol.

Intestinal antiseptic. Useful in typhoid. — *P. J.* '57, ii 83.

Dose — 8 to 10 grains = 0.52 to 0.65 gramme, three or four times a day.

Tests — Quinaphthol fuses at 185°C (365°F), when ignited with free access of air it leaves no weighable residue.

Sodium-Naphthol (Microcidin) readily soluble in Water, **Hydro-naphthol**, **Lactonaphthol (Lactol)**, and **Naphthol-Camphor** have also been introduced as possessing properties similar to those of Naphthol, **A-Oxy-naphthoic Acid** forms soluble salts with alkalis, which are antiseptics.

Not Official

NICKEL

A metal closely allied to Cobalt, with which it is generally associated in minerals. Commercially it is largely contaminated with Copper, Iron, and sometimes Cobalt. Alloyed with Copper and Zinc, it forms **German Silver**. Easily soluble in mineral acids, forming salts of a characteristic green colour.

NICCOLI BROMIDUM — Green, hygroscopic crystals, soluble in Water, Alcohol, and Ether.

Sedative. Recommended in epilepsy.

Dose — 1 to 5 grains = 0.06 to 0.32 gramme.

In solution, or in pills.

SYRUPUS NICCOLI BROMIDI — Granulated Nickel, 197 grains, Bromine, 377 grains, Water, 12 fl oz, digest them in a pint flask at a gentle heat until reaction ceases, filter, add Sugar, 24 oz, and sufficient Water to make 32 fl oz.

Each fl dram contains 5 grains of Nickel Bromide, which is an average dose
—*A J P* '86, 592

NICCOLI SULPHAS —Greenish-blue crystals, readily soluble in Water

Dose — $\frac{1}{2}$ to 1 grain two or three times a day in chlorosis, is best given on a full stomach, as otherwise it is apt to produce nausea. In somewhat larger doses it has also been given in locomotor ataxy.

The toxicology of **Nickel Carbonyl** —*B M J* '07, 11 687

Not Official

NITROGLYCERIN

Syn —GLYCERYL TRINITRATE, GLONONIN, TRINITRIN, TRINITRO-GLYCERIN

$C_3H_5(NO_3)_3$, eq 225 47

When pure it is a heavy, colourless, oily liquid. Explodes violently on percussion, and under some circumstances spontaneously.

A 10 p c solution in Alcohol is commercial, and is used in making the Tabellæ.

Solubility —Very slightly soluble in Water, readily in Alcohol (90 p c), mixes with Ether and with Chloroform.

Medicinal Properties —Chiefly given for angina pectoris associated with aortic disease, spasmodic asthma and the dyspnoea of acute bronchitis in hæmoptysis and in headache, neuralgia or hemicrania if associated with pain. It reduces arterial tension in chronic Bright's disease and acts as a diuretic and diminishes the albuminuria. Chemically, it is a nitrate, but its physiological action resembles the Nitrites. It is similar to that of Amyl Nitrite, but its action is slower and more prolonged.

Of service in migraine, especially when combined with Strychnine, but not for the attack itself, its service is obtained by continuous administration between the attacks (Gowers) —*B M J* '06, 11 1622

Preferred to Amyl Nitrite by some in hæmoptysis, because of its more lasting effect —*L* '08, 1 565

For hæmoptysis give two tabellæ in the 24 hours, breaking them up into small pieces, and letting the patient take a piece every hour —*P* '07, 1 335

In optic atrophy, *M A* '95, 261, in sciatica, in uræmic dyspnoea, in all forms of vomiting —*M P* 11 '95, 87, 445, 497, 520, *P* 11 140, in arterio-sclerosis, *T G* '93, 736, in warding off, and (hypodermically) during paroxysm of epilepsy, *B M J E* '93, 11 32, in gall-stone colic, *L* '00, 1 353

Dose — $\frac{1}{100}$ to $\frac{1}{50}$ grain = 0 0008 to 0 0013 gramme, the average dose being $\frac{1}{100}$ grain = 0 0008 gramme

Prescribing Notes —*The Solution may be given on Sugar, or in the form of Tablets, or diluted with Water*

Official Preparations —Liquor Trinitrini and Tabellæ Trinitrini

Not Official —Haustus Trinitrini, and Tabella Nitroglycerini Composita

Antidotes —Ergot, Atropine, Strychnine, cold applications to the head

Foreign Pharmacopœias —Official in Jap, Mex and Span.

Tests —Nitroglycerin has a sp. gr of 1 6 and, when pure, is colourless, but the commercial product generally has a yellow colour. It solidifies at 8° C (46° F) and is then very dangerous to handle. When smartly struck or compressed or when dropped on an iron plate heated to 257° C (494 6° F), it explodes with great violence. It is decomposed by Alcoholic Potassium Hydroxide Solution, yielding a mixture of Potassium Nitrate and Nitrite, Potassium Acetate and Formate. When treated with a solution of Ferrous Sulphate acidified with Hydrochloric Acid, gives the brown coloration characteristic of Nitrites and Nitrates.

Official Preparations**LIQUOR TRINITRINI** SOLUTION OF TRINITRIN *BP Syn* —
SOLUTION OF NITROGLYCERIN

Trinitroglycerin of commerce, 17½ grains, Alcohol (90 p c), q s to yield 4 fl oz

1 minim contains $\frac{1}{112}$ of a grain

Dose — $\frac{1}{2}$ to 2 minims = 0.03 to 0.12 c c

In severe cases of angina pectoris or asthma, the dose is sometimes increased

Foreign Pharmacopœias—Official in Dutch (Solutio Nitroglycerini), Dan, Jap, Mex and Span, 1 in 100, U S (Spiritus Glycerilis Nitratiss), 1 in 100. Not in the others

Tests—Solution of Nitroglycerin has a sp gr 0.830 to 0.836. The official gravity is 0.840. It is a clear, colourless liquid, possessing a neutral reaction towards Litmus paper. The presence of a due amount of Nitroglycerin is officially ensured by a test of which the following are the essential details—A measured quantity of 10 c c of the solution when mixed with 10 c c of Water yields a clear solution when cooled to 15.5° C (60° F), but a turbidity is produced in the mixture upon the further addition of 1 c c of Water, the Nitroglycerin separating out as an oily liquid when the mixture is still more largely diluted. If only 1 drop of this oily liquid be placed upon bibulous paper and sharply struck with a hammer a violent explosion results

TABELLÆ TRINITRINI TRINITRIN TABLETS *BP Syn* —
TABLETS OF NITROGLYCERIN

These tablets, made of chocolate, now weigh 5 grains instead of the 2½ grains in *BP* '85, but they contain as formerly $\frac{1}{100}$ of a grain = 0.0006 gramme of Trinitroglycerin

Dose — 1 or 2 tablets

Not Official

HAUSTUS TRINITRINI—Solution of Trinitrin, 1 minim Spirit of Chloroform, 5 minims, Tincture of Capsicum, 2 minims, Peppermint Water, to ½ oz — *Westminster*

TABELLA NITROGLYCERINI COMPOSITA—Nitroglycerin, $\frac{1}{100}$ grain, Menthol, $\frac{1}{80}$ grain, Capsicin, $\frac{1}{100}$ grain, Theobroma Paste, q s — *Westminster*

Not Official

NUCLEIN NUCLEOL

The nucleins are compounds of simple proteins with phosphorised bodies, and occur in Yeast, Milk, Yolk of Egg, Thyroid and Thymus glands, etc. Numerous varieties are supposed to exist. Nuclein is extracted by digestion with Pepsin and dilute Hydrochloric Acid, and purification of the residue by repeated solution and precipitation in dilute alkali and dilute acid respectively.

Nuclein is the best known chemical constituent of the nucleus of the white blood corpuscles

NUCLEIC OR NUCLEINIC ACID—A white or greyish white amorphous powder, slightly soluble in Water, insoluble in Alcohol (90 p c), and in Ether. It is readily soluble in solutions of Sodium or Potassium Hydroxides with the

formation of the so-called Nucleinates, and it is in the form of 5 p c aqueous solutions only that it is chiefly used medicinally

Medicinal Properties — A powerful germicide. Stated to possess nutritive properties, and to be useful in increasing the resisting power of the system to pathogenic germs. It has been employed in the treatment of tuberculosis, in anæmia, and in neurasthenia. It has also been employed in diphtheria, and puerperal and scarlet fevers.

1 p c solution of nucleinate of Sodium, in physiological solution, subcutaneously injected to moderate peritoneal inflammation after perforation in typhoid, and thus to lessen the risks of subsequent surgical interference. Within 3 days, 3 doses are given — *B M J* '07, 1 1515

Prescribing Notes — *It may be administered hypodermically in the form of a 5 p c solution, dose 17 minims = 1 c c, or by the mouth as a solution of its strength, in doses of 1 to 2 fl drms = 3.6 to 7.1 c c*

ARGENTI NUCLEINAS (Nargol) — A light brownish-yellow powder, containing about 10 p c Silver, soluble 1 in 4 of Water. Used as an injection, $\frac{1}{2}$ to 1 p c solution, in gonorrhœa — *B M J* '01, 1 1833, *L* '01, 1 1809, *M A* '02, 701

CUPRI NUCLEINAS (Cupiol) — A green, colourless, impalpable powder, soluble in Water. Its solution is stated not to coagulate albumen. Has been found in powder form in cases of trachoma. Also as a 5 to 10 p c solution in conjunctivitis. Is stated to be less irritating than the *U* '01, 1 729, 1809

FERRI NUCLEINAS (Triferrin Ferrinol) — A brown, amorphous, odourless powder, soluble in Water. Has been recommended in anæmia — *B M J E* '02, 1 104, 1 16, 104, *C D* '02, 1 580

Dose — 5 grains = 0.32 gramme

HYDRARGYRI NUCLEINAS (Mercuriol) — A pale yellowish-brown amorphous powder, soluble in Water, insoluble in Alcohol (90 p c). Its solution does not coagulate albumen. It has been used as a 2 p c injection in urethritis — *L* '00, 1 871. As an antiseptic in the form of a 2½ to 5 p c solution in the treatment of diseases of the nose and ear — *L* '00, 1 1726, *T G* '01, 92. Given with success in the treatment of syphilis, in average doses of 2 grains three times a day — *L* '01, 1 1039. In gonorrhœa as an injection in the form of a 2 p c solution — *T G* '01, 15. Has been found useful in combination with Chlorotone and Botic Acid in the treatment of various acute and chronic affections of the skin and mucous membrane — *T G* '01, 686

SODII NUCLEINAS — A white or greyish-white amorphous powder, soluble in Water. Employed medicinally, as above described, in the form of a 5 p c solution

NUX VOMICA.

NUX VOMICA

FR, NOIX VOMIQUE, GER, BRECHNUSS, ITAL, NOCE VOMICA,
SPAN, NUEZ VOMICA

The dried ripe Seeds of *Strychnos Nux-vomica*, L

Imported from India, Ceylon, and Cochin China

The chief source of Strychnine and Brucine

The total alkaloids have been found to vary between 1.25 and 3.9 p c (some Ceylon Seeds gave 5.3 p c), but the value of total alkaloids as a medicinal standard is considerably reduced by the fact that the ratio of Strychnine to Brucine may vary as much as 3 to 1 and 1 to 2. The official galenical preparations of *Nux Vomica* are

standardised to a definite percentage of Strychnine, but the *B P* does not state what amount of Strychnine should be present in the Seeds. The *U S P* requires that they shall yield not less than 1.25 pc of Strychnine, the *P G* not less than 2.5 pc of total alkaloids as calculated from the result of the official volumetric process, using a factor based on equimolecular proportions of Strychnine and Brucine. The *Brussels Conference* has agreed upon a standard of 2.5 pc of total alkaloids. The *Fr Codex* (1908) has adopted the recommendation of the *Brussels Conference*, and requires that the dried powdered seeds should yield not less than 2 nor more than 3 pc of total alkaloids. From 1 to 1.25 pc of Strychnine has been suggested (*Y B P* '03, 252) as a suitable standard for inclusion in the next *B P*, the dual standard of percentage of total alkaloids and percentage of Strychnine not being advocated. The establishment of a Strychnine standard upon a dual basis by calculating the Brucine into terms of Strychnine has been suggested, *Y B P* '06, 237.

Medicinal Properties—An excellent gastric and general tonic. Recommended in atonic dyspepsia, in general debility, and in convalescence. It stimulates peristalsis, and therefore is a frequent and valuable ingredient in medicines for chronic constipation. It is also a cardiac and respiratory stimulant. Useful in paralysis of reflex origin, in peripheral paralysis due to alcohol, lead, tobacco, or to diphtheria, in all chronic paralytic affections, except those in which there is organic lesion of nerve-centres or inflammation of brain or spinal cord. See also STRYCHNINA.

A report on eight cases of chronic pulmonary tuberculosis, treated with a 'simple mixture of nux vomica, gentian and acid,' compared with similar cases treated with 'Malt and Oil,' results compare most favourably.—*L* '03, ii 1016.

Dose—In powder 1 to 4 grains = 0.06 to 0.26 gramme.

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.2 gramme.

Prescribing Notes— $\frac{1}{15}$ grain Strychnine is contained in $1\frac{1}{2}$ grains of *Extract*, $5\frac{1}{2}$ minims of *Fluid Extract*, 33 minims of *Tincture*.

Official Preparations—Of the seeds, *Extractum Nucis Vomice Liquidum* and *Strychnina*, of the *Liquid Extract*, *Extractum Nucis Vomice* and *Tinctura Nucis Vomice*.

Not Official—Brucine.

Antidotes—Emetic of Zinc Sulphate, Mustard, or Ipecacuanha, or hypodermic injection of Apomorphine, Animal Charcoal, Potassium Bromide or Chloral, Amyl Nitrite inhalations, Chloroform or Ether to relax the muscles, hypodermic injection of Curare.—*Murrell*.

Foreign Pharmacopœias—Official in Austri, Belg., Dutch, Ger., Jap., Russ. and Swiss, *Semen Strychni*, Dan., Fr. (*Noix Vomique*), Hung., Ital. (*Noce Vomica*), Mex. and Span. (*Nuez Vomica*), Norw., Port. (*Noz Vomica*), Swed. and U S.

Descriptive Notes.—*Nux Vomica* Seeds are imported from Ceylon, Bombay, Cochin, Madras, and Calcutta. They vary in size and in alkaloidal content, the largest usually yielding most alkaloid, the Ceylon and Bombay Seeds are richer than those imported from Madras and Cochin. The Seeds vary from $\frac{3}{4}$ to 1 in (19 to 25 mm.)

in diameter and from $\frac{1}{8}$ to $\frac{1}{4}$ in (3 to 6 mm) in thickness. They are circular, nearly flat or somewhat plano-convex, and occasionally irregularly bent, greyish-green in colour, with a satiny lustre from the appressed hairs, the margin rounded or in some kinds acute with a protuberance at the edge indicating the position of the radicle, internally consisting of hard, tough, and horny albumen, in the centre of which is found an embryo with thin, leafy, cordate, palmately-veined cotyledons. The taste is intensely and persistently bitter. *BP* gives the same measures as above for the Seeds, *USP* diameter 15 to 30 mm, thickness 3 to 5 mm. The microscopical characters of the powder are the non-porous thick-walled endosperm cells (*PG*), containing fixed Oil and aleurone grains (*USP*), the hair bases with linear specially formed pits and the cylindrical fragments of the upper part of the hairs, which have a striated appearance.

Tests—Nux Vomica Seeds may be assayed for their percentage content of Strychnine by any one of several excellent processes. That of the *USP* is essentially as follows—A weighed quantity of 20 grammes of the Seeds in No. 60 powder is introduced into an Erlenmeyer flask and is first macerated for 1 hour, with frequent intervals of shaking, with 200 cc of a mixture composed of 137.5 cc of Ether, 44 cc of Chloroform and 18.5 cc of Alcohol (94.9 pc) and 5 cc of Ammonia Solution and subsequently allowed to stand for 12 hours. A measured quantity of 100 cc of the mixture is decanted into a second separator, the residue is washed with a little Chloroform and the washings are added to the first. The alkaloids are then extracted from the chloroformic solution by agitation with 15 cc of Normal Volumetric Sulphuric Acid Solution, care being taken to avoid the formation of emulsion during the shaking, after complete separation the lower acid layer is separated, the Ether-chloroform solution is separated, washed with 2 successive quantities each of 5 and 3 cc of Normal Volumetric Sulphuric Acid Solution. The acid liquids in each case are separated as previously and mixed with the main acid quantity. The complete extraction of the alkaloids from the Ether-Chloroform liquid is ensured by testing a drop of the acid liquid with Mercuric Potassium Iodide (Mayer's) Solution, and if a precipitate is produced the shaking is repeated with a further quantity of 5 cc of Normal Volumetric Sulphuric Acid Solution, the acid solutions are mixed and sufficient Ammonia Solution added to render the solution alkaline, and the liberated alkaloids are extracted by thoroughly shaking first with 25 cc of Chloroform and subsequently repeating the extraction with two successive portions, each of 15 cc of Chloroform. The chloroformic solution is in each case separated, transferred to a tared flask, the mixed chloroformic solutions evaporated to dryness on a water-bath and the residue dissolved in 15 cc of 3 pc Sulphuric Acid by warming it on the water-bath. The solution is allowed to cool and 3 cc of a cooled mixture of equal volumes of Nitric Acid (sp. gr. 1.42) and Distilled Water added, the liquid, after it has been rotated a few times, is set aside for exactly 10 minutes, with 3 intervals of

gentle rotation, the liquid is transferred to a separator containing 25 c c of a 10 p c w/v Sodium Hydroxide Solution, the flask being washed out with 3 successive small quantities of Water and the washings added to the main quantity of liquid, which, if not turbid, is mixed with a further measured quantity of 2 c c of the Sodium Hydroxide Solution. The liberated alkaloids are then extracted by well rotating the mixture for a few minutes with 20 c c of Chloroform, the extraction being repeated with 2 successive quantities of 20 c c of Chloroform, the Chloroform solution is separated in each case, filtered through a small filter paper, previously moistened with Chloroform, into a tared flask, the filter and funnel washed with 5 c c of Chloroform and the mixed Chloroform solutions evaporated very carefully to dryness on a water-bath. The alkaloidal residue is dissolved in 6 c c of Tenth normal Volumetric Sulphuric Acid Solution, 5 drops of Iodoosin Test Solution are added, about 80 c c of Water and 20 c c of Ether, the excess of acid is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution. The number of c c of Fiftieth-normal Potassium Hydroxide Solution is divided by 5, the quotient subtracted from 6, the difference multiplied first by 0.0332 and then by 10 yields the percentage of Strychnine present in the sample. The above process for the separation of Strychnine and Brucine is that originally suggested by Gordin, but, as originally introduced into the *U S P*, it was modified in two essential particulars, such modifications having been vigorously protested against by the author of the process, who considered the modifications quite unwarranted. The method originally suggested by Gordin (*Proc Amer Pharm Assoc* '02, 341) states, first, that the mixed alkaloids might be dissolved in 15 c c of a 3 p c Sulphuric Acid Solution by the aid of the water-bath heat, and after the solution is cooled to the ordinary temperature 3 c c of a specially prepared and cooled mixture of equal parts of strong Nitric Acid (sp gr 1.420) are to be added. Secondly, the mixed Chloroform solutions containing the residual Strychnine are directed to be mixed with 2 or 3 c c of pure Amyl Alcohol [b p 128° to 132° C (262.4° to 269.6° F)], previous to evaporation to dryness. Nitric Acid of a sp gr of 1.40 does not affect the oxidation of the Brucine to the extent that an acid of a sp gr of 1.42, and requires the addition of a small quantity of Sodium Nitrite to start the reaction. The insufficiency of the original *U S P* official Nitric Acid (sp gr 1.40) had been also pointed out (*Proc Amer Pharm Assoc* '07, 55, 781), however carefully the evaporation of the Chloroform solution is conducted, without the addition of the Amyl Alcohol there is a considerable liability to a loss due to decrepitation, but with the addition of 2 or 3 c c of pure Amyl Alcohol no decrepitation takes place. It will be noted that the *U S P* omits the use of Amyl Alcohol. Farr and Wright (*Y B P* '06, 226) have experimented with the *U S P* Nitric Acid process for the determination of Strychnine, they appear to be of the opinion that, notwithstanding the condemnation of the process as contained therein, that process, with slight modifications in the working details, gives perfectly satisfactory results, and that subsequent work has thoroughly established its

reliability The exact details of the process as they have applied it are as follows —The total alkaloids obtained in the usual way from 5 cc of the liquid extract or 25 cc of the Tincture are dissolved by the heat of a water-bath in 15 cc of 3 p c Sulphuric Acid Solution, the temperature of the solution adjusted to 50° C (122° F), 3 cc of a mixture of equal volumes of Nitric Acid (sp gr 1.42) and Water added and the mixture set aside for 10 minutes, it is transferred to a separator and shaken with Chloroform, the Chloroform solutions run into a tared dish containing 3 cc of Amyl Alcohol. It will be noticed that the above modification by Farr and Wright of the *USP* process rectifies the identical objections which were urged against the strictly *USP* process, namely, the gravity of the Nitric Acid Solution used for the oxidation of the Brucine and the addition of Amyl Alcohol to prevent loss by decrepitation of the Strychnine residue. The original process of the *USP* recommended the use of Nitric Acid of specific gravity 1.40, but the list of alterations and corrections (1907) has altered the specific gravity of the acid to 1.42.

Another modification of the *USP* method of separating Brucine and Strychnine is described (*AJP* '07, 6). The alkaloidal residue is dissolved in 15 cc of 3 p c Sulphuric Acid, to the solution is added 3 cc of a mixture of equal volumes of Nitric Acid (sp gr 1.4) and Distilled Water. Then add 1 cc of a 5 p c Solution of Sodium Nitrite in Water and, after rotating the liquid a few times, set it aside for exactly 30 minutes, stirring it gently 3 times during the interval, the solution is then made alkaline and shaken out with Chloroform in the usual way.

The influence of Nitrous Acid in the oxidation of Brucine by Nitric Acid has been closely studied by Reynolds and Sutcliffe (*JSCI* '06, 512), and they conclude that Stoeder's and Gordin's addition led to slightly more accurate results than Keller's original process, of the two, that of Gordin should have the preference as it is more expeditious. The short Nitric Acid process is generally capable of accurate results, if the following points are attended to —(1) For the amount of total alkaloid up to 0.4 of a gramme, the reacting solution should contain at least 7 p c of Nitric Acid. (2) The reaction should be stopped after 10 minutes, when the Brucine is entirely oxidised. (3) The temperature should not exceed 25° C (77° F). (4) Excess of Potassium or Sodium Hydroxide should be used to liberate the Strychnine, and not Sodium Carbonate or Ammonia Solution. (5) The Nitric Acid used should be added in the form of sp gr 1.42 and not more diluted, otherwise it may be necessary to add a trace of Nitrate to start the reaction.

The *PG* process is a volumetric one. A quantity of 15 grammes of the Seeds dried at 100° C and reduced to a middling fine powder is shaken with 100 grammes of Ether and 50 grammes of Chloroform, and then mixed with 10 cc of a solution of 2 parts by weight of Sodium Hydroxide Solution (15 p c) and 1 part by weight of Water, and the residue is allowed to stand for 3 hours with intervals of vigorous shaking. A measured quantity of 15 c.c. or

a sufficient quantity of Water to cause the powdered Nux Vomica to agglomerate and the supernatant Chloroform-Ether solution to separate completely clear, is then added. After standing for an hour 100 grammes of the clear Ether-Chloroform solution are filtered through a dry, well-covered filter into a flask, and about half of the liquid distilled, the remaining Chloroform Ether solution is transferred to a separator, the flask being washed out with 3 successive quantities, each of 5 c c of a mixture of 3 parts by weight of Ether, and 1 part by weight of Chloroform. The alkaloids are extracted from the mixed liquids by agitation with 10 c c of Tenth normal Volumetric Hydrochloric Acid Solution. After complete separation of the liquids sufficient Ether is added to cause the Chloroform Ether solution to float on the top of the acid liquid, the latter is filtered through a small filter previously moistened with Water into a flask of 100 c c capacity. The Chloroform-Ether solution is shaken with 3 successive quantities, each of 10 c c of Water, the aqueous liquids being filtered through the same filter, the filter is washed with Water and the mixed filtrate and washings diluted with Water to 100 c c. A measured quantity of 50 c c of this solution is removed, introduced into a flask of white glass of about 200 c c capacity, 50 c c of Water and sufficient Ether to form a layer of 1 cm added, and the mixture titrated with Hundredth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosm Solution as an indicator of neutrality, not more than 15 c c of Hundredth normal Volumetric Potassium Hydroxide Solution shall be necessary to neutralise the excess of Tenth normal Hydrochloric Acid, the number of c c of Hundredth-normal Volumetric Hydroxide Solution used divided by 10, the quotient multiplied first by 2, then subtracted from 10 and the difference multiplied by 0.00364 (the mean molecular equivalents of Strychnine and Brucine), then by 10 gives the percentage of total alkaloid present in the Seeds.

Preparations

EXTRACTUM NUCIS VOMICÆ EXTRACT OF NUX VOMICA

Prepared from Liquid Extract of Nux Vomica, and readjusted by means of Milk Sugar to contain 5 p c of Strychnine.

The *BP* extract is prepared by the evaporation of the liquid extract, which is officially required to contain 1.5 p c w/v of Strychnine. The solid extract is officially required to contain 5 p c of Strychnine.

The *USP* extract is prepared direct from the powdered Seeds, the menstruum being a mixture of Acetic Acid and Water, it is required to contain 5 p c of Strychnine. The *PG* extract is also prepared from the powdered Seeds, using the menstruum Alcohol (68 to 69 p c), and is required to contain not less than 17.5 p c w/w of mixed alkaloids. The Extract official in the *Fd Codex* (1908) is prepared from the Nux Vomica Seeds in No. 22 powder, the menstruum being Alcohol (70 p c). It is required to contain exactly 16 p c of total alkaloids in conformity with the recommendation of the *Brussels Conference*.

Dose.— $\frac{1}{4}$ to 1 grain = 0.016 to 0.065 gramme

Ph Ger maximum single dose, 0.05 gramme, maximum daily dose, 0.10 gramme

Often prescribed with Aloes and Ipecacuanha

This Extract is intended to be about two-thirds the strength of that in *B P* 1885. *Ph Ger* Extractum Strychni is standardised to contain 17.5 p.c. of total alkaloids, and is therefore rather stronger than *B P* 1885

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Jap., Russ., Span. and Swiss, use 68 to 70 p.c. Alcohol, Ital. and Mex., 80 p.c., Norw. and Swed., 65 p.c., Port., 90 p.c., U.S. extract with Acetic Acid and Water, and subsequently add Alcohol (95 p.c.) All from the Seeds. Austr., Belg., Dutch, Fr., Span. and Swiss adopt the International Standard, 16 p.c. of alkaloids, Dan. and Swed., 15 to 17 p.c., Ital., 10 p.c., Mex., 15 p.c., Russ., 15 p.c., U.S., 5 p.c. of Strychnine. Ger. and Jap., 17.5 p.c.

Tests.—No official method is given for the determination of the Strychnine in the *B P* extract, it being apparently assumed that the liquid extract used in its preparation contains the necessary proportion of alkaloid

The *U S P* dissolves in a beaker a weighed quantity of 2 grammes of the extract in 25 cc of a mixture of 16 cc of Ether, 5 cc of Chloroform, and 4 cc of Ammonia Solution, transferring it when dissolved to a separator and washing the beaker with a little Chloroform, transferring the washings also to the separator. The alkaloids are extracted by agitating the mixture for a few minutes, the aqueous layer is transferred to a second separator, the Ether solution and the first separator being washed with a little Water and the washings separated and added to the second separator. The alkaloids remaining in the aqueous liquid are extracted by agitation with 2 successive portions each of 15 and 10 cc of Chloroform, the Chloroform solutions are separated and added to the Ether solution contained in the first separator. The complete extraction of the alkaloids from the aqueous liquid is ensured by removing a few drops, rendering them acid and testing with Mercuric Potassium Iodide (Mayer's) Solution, and if a reaction is obtained repeating the shaking with a fresh quantity of 10 cc of Chloroform. The alkaloids are extracted from the mixed Ether-Chloroform solutions contained in the first separator by agitation with 3 successive quantities each of 15, 10, and 10 cc of 3 p.c. Sulphuric Acid Solution, the acid layer being in each case separated, mixed and transferred to another separator, sufficient Ammonia Solution to render the mixture alkaline is added and the liberated alkaloids are extracted by agitation with 3 successive quantities of 15 cc, 10 cc, and 10 cc of Chloroform, the Chloroform solutions are separated in each case, transferred to a beaker, and evaporated on a water-bath. The alkaloidal residue is dissolved in 15 cc of a 3 p.c. Sulphuric Acid Solution whilst still on the water-bath, removed and allowed to cool. A measured quantity of 3 cc of cooled mixture of equal volumes of Nitric Acid (sp. gr. 1.42) and Water are added, the liquid rotated a few times, set aside for exactly 10 minutes, during which it is gently stirred 3 times. The red liquid is transferred to a separator containing 25 cc of a 10 p.c. w/v Sodium Hydroxide Solution, the beaker washed with 3

successive very small amounts of Water and the washings transferred to the separator, a further quantity of 2 c c of the Sodium Hydroxide Solution being added, should the liquid be not quite turbid. The liberated alkaloids are extracted by agitation with 20 c c of Chloroform, adopting a rotatory motion, the complete extraction of the alkaloids being ensured by shaking with 2 further successive quantities of 10 c c of Chloroform, the same rotatory method being adopted in the shaking, the Chloroform solutions are separated in each case, filtered through a small filter previously wetted with Chloroform into a tared flask, the filter and funnel are washed with 5 c c of Chloroform. The mixed Chloroform liquids evaporated very carefully to dryness on a water bath, the alkaloidal residue is dissolved in 10 c c of Tenth-normal Volumetric Sulphuric Acid Solution, about 90 c c of Distilled Water and 20 c c of Ether and about 5 drops of Iodeosin Solution are added and the excess of Volumetric Acid Solution is titrated with Fiftieth normal Volumetric Potassium Hydroxide Solution, the number of c c used is divided by 5, the quotient is subtracted from 10, the difference is multiplied first by 0.0332 and then by 50, which yields the percentage of Strychnine present in the extract.

The *P G* dissolves a weighed quantity of 1 gramme of the extract in 5 grammes of Water and 5 grammes of Absolute Alcohol, and adds to this solution 50 grammes of Ether and 20 grammes of Chloroform, and after vigorous shaking adds 10 c c of a 1 in 3 solution of Sodium Carbonate and allows the mixture to stand for an hour with frequent intervals of vigorous agitation. A weighed quantity of 50 grammes of the clear Chloroform solution is filtered through a dry, well-covered filter into a flask, and about half of the liquid is distilled, the remaining Chloroform solution is introduced into a separator, the flask is washed with 3 successive quantities each of 5 c c of a mixture of 3 parts by weight of Ether, and 1 part by weight of Chloroform, and the alkaloids are extracted from the mixed liquids by shaking with 50 c c of Hundredth normal Volumetric Hydrochloric Acid Solution. When the clear liquids have completely separated, and after the addition of sufficient Ether to cause the Chloroform-Ether solution to float on the acid liquid, the latter is separated, filtered through a small filter previously moistened with Water into a stoppered flask of white glass of a capacity of about 200 c c. The Chloroform-Ether solution is washed with 3 successive quantities each of 10 c c of Water, the washings filtered through the same filter, the filter washed with Water, and the mixed filtrate and washings diluted with Water to about 100 c c. After the addition of sufficient Ether to form a layer of 1 cm, sufficient Hundredth-normal Volumetric Potassium Hydroxide Solution is added to neutralise the excess of the Volumetric Acid Solution, Iodeosin Solution being employed as an indicator of neutrality. The number of c c of hundredth normal alkali solution subtracted from 50, the difference multiplied by 0.00364 (the mean molecular equivalent of Strychnine and Brucine) and the product multiplied by 100, and this product divided by 0.711 yields the percentage of total alkaloids present in the extract.

EXTRACTUM NUCIS VOMICÆ LIQUIDUM LIQUID EXTRACT
OF NUX VOMICA

A fluid prepared by percolation with Alcohol (70 p c) and standardised to contain 1·5 grammes of Strychnine in 100 c c

The liquid extract is officially required to contain 1·5 p c w/v of Strychnine. The *USP* Fluid Extract is required to contain 1 p c w/v of Strychnine, neither the *PG* nor the *Fr. Codex* (1908) contains a Fluid Extract of Nux Vomica.

Dose.—1 to 3 minims = 0·06 to 0·18 gramme

Foreign Pharmacopœias—Official in Mex and US. The *USP* extracts with a mixture of Alcohol (95 p c) 3, Water 1, to which Acetic Acid has been added, it is standardised so that each 100 c c of finished Fluid Extract shall contain 1 gramme of Strychnine.

The *Brussels Conference* agreed to prepare the Extract by means of Alcohol (70 p c) and to an alkaloidal strength of 16 p c.

Tests.—Fluid Extract of Nux Vomica has a sp gr of 0·945 to 0·965, it contains from 9 to 12 p c of total solids and about 58 p c w/v of Absolute Alcohol. The proportion of total solids may amount to as much as 20 p c, but will largely depend upon the Strychnine content of the Seeds used in the liquid extract. It has been pointed out (*YBP* '06, 236) is evident, in fixing the official standard at 1·5 p c, somebody has blundered, for it is palpably impossible to produce from a drug which rarely contains as much as 1·5 p c of Strychnine, a 1 in 1 preparation standardised to contain that amount.

The *BP* method of determination, which has been very severely and adversely criticised, is essentially as follows.—A measured quantity of 10 c c of the liquid extract is evaporated to the consistency of a thick syrup by heating on a water-bath, and the resulting extract is dissolved in 20 c c of Water, transferred to a separator and the solution mixed with a solution of 5 grammes of Sodium Carbonate in 25 c c of Water. The liberated alkaloids are extracted by agitation with 3 successive quantities each of 10 c c of Chloroform, the Chloroform solutions being in each instance separated and transferred to a second separator. Unless the fat has been previously extracted from the Seeds a considerable quantity of fatty matter is present, which gives rise to the formation of troublesome emulsions at this stage of the process, the Chloroform obstinately emulsifies and refusing to separate. The alkaloids are in turn extracted from the mixed chloroformic liquids by agitation with 3 successive quantities each of a third part of a mixture of 6 c c of diluted Sulphuric Acid with 25 c c of Water. The acid solutions are in each instance separated, mixed, diluted with Water to 175 c c and transferred to a stoppered bottle in which after being made up to 200 c c with Potassium Ferrocyanide Solution they are well and frequently shaken for thirty minutes. The separation of the mixed alkaloids is based on the insolubility of the Strychnine as compared with that of the Brucine Ferrocyanide, an observation due to Beckurts, and subsequently utilised by Dunstan and Sholl (*YBP*, 1883, 469), for the determination of

Strychnine in the presence of Brucine Unless strict attention be paid to the temperature at which the precipitation is carried out and that the solution be well agitated during the addition of the Ferrocyanide, Brucine will be also precipitated with Strychnine. The mixture is allowed to stand for 6 hours to allow complete precipitation, the supernatant fluid is decanted, filtered through a small filter, the precipitate transferred to the same filter, the precipitate remaining in the bottle is also transferred to the filter by washing it out with some of the clear filtrate and then with Water containing one-fortieth of its volume of diluted Sulphuric Acid, and the precipitate is washed until the washings are free from bitterness. If this stage of the process be strictly followed considerable loss of alkaloid will result, as the Strychnine Ferrocyanide is not completely insoluble in this menstruum. The precipitate is then transferred to a separator, mixed with 5 c c of Ammonia Solution, and well shaken, and the liberated alkaloids are extracted by agitation with 2 successive quantities each of 15 c c of Chloroform, the Chloroform solutions in each instance being separated, transferred to a tared flask and the mixed chloroformic liquids evaporated on a water-bath, the residue being dried for 1 hour at a water bath temperature, or preferably till constant in weight, cooled and weighed. If the final Chloroform solutions are evaporated, as officially directed, in a counterpoised dish, there is considerable liability to loss by decrepitation, which may be avoided by the addition of a few c c of Amyl Alcohol to the chloroformic liquids previous to evaporation. The calculated yield of alkaloids from the 10 c c of liquid extract directed by the *B P* to be employed in the test is 0.32 of a gramme, and the process upon which the separation was founded directs the employment of any quantity of liquid yielding not more than 0.2 of a gramme of mixed alkaloids. Several useful suggestions have been made with a view of overcoming the defects of the Pharmacopœia process. Farr and Wright have suggested (*Y B P* '00, 450) the following modifications: that the volume of liquid taken should not exceed 5 c c of the liquid extract or 30 c c of Tincture, that 200 c c of Water at a stated temperature, preferably 38° C (100.4° F) should be employed and a correction made for the Strychnine dissolved, and that in carrying out the process the Pharmacopœia instructions as to a simple agitation without stirring and as to the length of time allowed for precipitation of the Strychnine are to be strictly observed, as success depends altogether upon the conditions under which the process is carried out. Bud has suggested the addition of 2 c c of pure Amyl Alcohol to the final Chloroform solution of the alkaloid before evaporation: it prevents decrepitation of the Strychnine when the residue dries. He has also suggested the removal of the fat by a preliminary shaking out with Chloroform in acid solution before starting the assay, the traces of alkaloid dissolved by the Chloroform solution of the fat being recovered by shaking the Chloroform solution again with acid. Naylor (*Y B P* '05, 364) is of opinion that the difficulties attending that part of the official process which refers to the separation of the

Brucine. Even assuming the adoption of the minute precautions proposed by Farr and Wright, cannot confidently be affirmed to have been surmounted, and it is imperative that attention to details of an unusually exacting character be carefully observed if results claiming to be concordant are to be obtained. Later results have shown that a more expeditious, easier, and certainly a more accurate method of separating Brucine from Strychnine consists in oxidising it with Nitric Acid in the presence of Sulphuric Acid. The process was worked out by Gordin and is given *in extenso* in the *Proceedings of the American Pharmaceutical Association* 1902, vol 1, p 336. The residue of total alkaloids obtained in the assay of Nux Vomica or its preparations is dissolved in 15 cc of 3 p.c. Sulphuric Acid Solution by the aid of a water-bath heat, the solution is cooled to ordinary temperature and 3 cc of a previously prepared and cooled mixture of equal parts of strong Nitric Acid (sp gr 1.42) and Water added to the alkaloidal solution, the liquid is set aside for exactly 10 minutes, shaking it gently 3 or 4 times during this time. The red liquid is transferred to a separator containing 20 to 25 cc of a 10 p.c. w/v Sodium Hydroxide Solution, and the vessel in which the digestion of the alkaloids has taken place is washed 3 or 4 times with very small amounts of Water, the washings being added to the contents of the separator. In the event of the liquid not being turbid a further addition of 1 or 2 cc of the Sodium Hydroxide Solution should be made, the liberated alkaloids are shaken out with 3 successive quantities of 20 cc, 10 cc and 10 cc of Chloroform. The chloroformic liquids are separated, filtered through a small filter paper previously moistened with Chloroform into a tared flask, the filter washed with Chloroform. To the mixed Chloroform solutions are added 2 or 3 cc of pure Amyl Alcohol distilling between 128° and 132° C (262.4° to 269.6° F), the mixed solutions are evaporated to dryness, the residue dried for about 2 hours at a temperature of 135° to 140° C (275° to 284° F), and when cool. If the fat be removed, as suggested above by Bird's process, the mixed alkaloids can be separated by the above process, the combination of the two processes yielding results of a satisfactory and concordant nature.

The U.S.P. method of determining the Strychnine in the Fluid Extract is as follows.—A measured quantity of 10 cc of the Fluid Extract is transferred to a porcelain evaporating basin and evaporated on a water-bath to dryness, the residue whilst warm being dissolved in a mixture of 16 cc of Ether, 5 cc of Chloroform, and 4 cc. of Ammonia Water, the solution being transferred to a separator, the dish rinsed with a little Chloroform and the rinsings added to the separator which is carefully shaken for a few minutes. After the liquids have separated the aqueous layer is removed to another separator, the Ether-Chloroform liquid is washed with a little Water and the washings added to the second separator, and the aqueous liquid in the second separator is shaken with 2 successive quantities each of 15 and 10 cc of Chloroform, which are added to the Chloroform solution in the first separator. Complete extraction

of the alkaloids is ensured by acidifying a few drops of the aqueous liquid remaining after Chloroform extraction and testing with Mercuric Potassium Iodide (Mayer's) Solution, if a reaction is obtained, a further shaking with 10 cc of Chloroform is carried out. The alkaloids are extracted from the Ether Chloroform mixed with the Chloroform liquids in the first separator by shaking with 3 successive quantities each of 25, 10 and 10 cc of Normal Volumetric Sulphuric Acid Solution. The acid solutions are separated in each case, transferred to a separator, and the mixed acid solutions rendered alkaline with sufficient Ammonia Solution, and the liberated alkaloids shaken out with 3 successive quantities each of 25, 10 and 10 cc of Chloroform. The Chloroform solutions are separated in each case, transferred to a beaker, and the mixed chloroformic liquids evaporated to dryness on a water-bath. The alkaloidal residue is dissolved in 15 cc of 3 p.c. Sulphuric Acid Solution by the heat of the water bath, allowed to cool, mixed with 3 cc of a cooled mixture of equal volumes of Nitric Acid (sp. gr. 1.42) and Distilled Water, and after rotating the liquid a few times, set aside for exactly 10 minutes, during which interval it is gently stirred on 3 successive occasions. The red liquid is transferred to a separator containing 25 cc of 10 p.c. w/v Sodium Hydroxide Solution, the beaker washed with 3 successive very small quantities of Water, and the washings transferred to the separator, a further quantity of 2 cc of the Sodium Hydroxide Solution is added in the event of the liquid not becoming turbid. The liberated alkaloids are extracted by shaking well (adopting a rotatory motion) with 3 successive quantities of 20, 10 and 10 cc of Chloroform, the Chloroform solutions in each case being separated, filtered through a small filter previously moistened with Chloroform, the filter and funnel washed with 5 cc of Chloroform, the mixed Chloroform solutions and washings carefully evaporated by means of the water-bath to avoid decrepitation, and the alkaloidal residue is dissolved in 10 cc of Tenth-normal Volumetric Sulphuric Acid Solution, about 80 cc of Water and 20 cc of Ether. The excess of acid is titrated with Fiftieth normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosin T.S. as an indicator of neutrality. The number of cc of Fiftieth normal Volumetric Potassium Hydroxide Solution required divided by 5, the quotient subtracted from 10, the difference multiplied first by 0.0332 and then by 10 yields the percentage of Strychnine present in the Fluid Extract.

TINCTURA NUCIS VOMICÆ TINCTURE OF NUX. VOMICA.
N O Syn — TINCtura STRYCHNI

Liquid Extract of Nux. Vomica, 2, Distilled Water, 3, Alcohol (90 p.c.), q.s. to yield 12

It is about twice the strength of the *B.P.* 1885 Tincture

The *B.P.* Tincture of Nux. Vomica is required to contain not less than 0.24 p.c. w/v nor more than 0.26 p.c. w/v of Strychnine. The *U.S.P.* Tincture is required to contain 0.1 p.c. w/v of Strychnine. The *P.G.* Tincture is required to contain not less than 0.25 p.c. w/v of mixed alkaloids.

The *BP* Tincture is prepared from the standardized liquid extract. The *USP* Tincture is prepared from the standardized extract. The *PG* Tincture is prepared from the powdered Seeds. The Tincture official in the *Ph Codex* (1908) is prepared in accordance with the recommendations of the *Brussels Conference*, namely, from the extract, using Alcohol (70 p c), and is required to contain 0.25 p c of total alkaloids.

Dose.—5 to 15 minims = 0.3 to 0.9 c c

Ph Ger maximum single dose, 1.0 gramme, maximum daily dose, 2.0 grammes.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Ger, Ital, Jap, Norw, Russ, Span, Swed and Swiss, 1 in 10, Hung, Mex and Port, 1 in 5, all prepared from the seeds. *US*, 1 Extract in 50. All by weight, except *US*.

Tests.—Tincture of *Nux Vomica* should have a sp gr of 0.890 to 0.915. It contains from 2 to 3 p c w/v of total solids and about 64 p c w/v of Absolute Alcohol. It is officially required to yield the percentage of Strychnine shown above. The *BP* method of determination is similar to that adopted for the assay of the liquid extract, 100 c c of the Tincture are evaporated to the consistency of a thick extract and the process continued as described under the Liquid Extract. The same comments as there appear are naturally applicable to the adaptation of the process to the Tincture.

The *USP* method is to evaporate a measured quantity of 100 c c of the Tincture to dryness on the water-bath, and to determine the amount of Strychnine present by the method of assay as given under *Extractum Nucis Vomice*. The final multiplication by 50 must in this instance be omitted, as the result will represent the percentage w/v of Strychnine present in the Tincture. The remarks regarding the Nitric Acid process for the separation of Brucine and Strychnine apply here. The German Pharmacopœia evaporates a weighed quantity of 50 grammes of the Tincture in a tared dish to the weight of 10 grammes and transfers the residue to a stoppered vessel with 5 grammes of Absolute Alcohol, the mixture is shaken with 50 grammes of Ether and 20 grammes of Chloroform, 10 c c of a 1 in 3 Sodium Carbonate Solution (which has been used to transfer the last traces of the residue left in the evaporating basin to the stoppered vessel) added, and the mixture allowed to stand for 1 hour with frequent intervals of vigorous shaking. A weighed quantity of 50 grammes of the clear Chloroform-Ether solution is filtered through a dry, well-covered filter into a flask, and about half the liquid distilled, the remaining Chloroform-Ether solution is introduced into a separator, the flask is washed with 3 successive quantities of 5 c c of a mixture of 3 parts by weight of Ether, and 1 part by weight of Chloroform, and the alkaloids are removed from the mixed fluids by shaking thoroughly with 40 c c of Hundredth-normal Volumetric Hydrochloric Acid Solution. After complete separation sufficient Ether is added to cause the Chloroform-Ether solution to float on the surface of the acid liquid. The latter is filtered through a small filter previously moistened with Water into a stoppered flask of about

200 c c capacity, the Chloroform-Ether solution is washed with 3 successive quantities each of 10 c c of Water and the washings filtered through the same filter, which is finally washed with Water, and the mixed filtrate and washings are diluted with Water to about 100 c c. After the addition of sufficient Ether to form a layer of about 1 cm, the excess of volumetric acid is titrated with Hundredth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosin Solution as an indicator of neutrality, the mixture being well shaken after each addition. It is required that not more than 17 c c of the Hundredth-normal Solution shall be necessary. The number of c c of Hundredth-normal Volumetric Potassium Hydroxide Solution required is subtracted from 40, the difference multiplied by 0.00364 (the mean molecular equivalents of Strychnine and Brucine), the product multiplied by 100 and divided by 33, yields the percentage w/w of total alkaloids present in the Tincture.

STRYCHNINE —*See* STRYCHNINA

Not Official

BRUCINE ($C_{23}H_{27}N_2O_4 \cdot 4H_2O$, eq 462.85) —Colourless, transparent, monoclinic crystals, containing about 15 p c of Water. Its salts are bitter, and most of them crystallisable.

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with air, as the crystals quickly effloresce when exposed to dry air.

Solubility —But slightly soluble in Water, 1 in 20 of Alcohol (90 p c), 1 in 2 of Chloroform, with separation of the combined Water.

Brucine resembles Strychnine in its physiological action, but is weaker.

Dose — $\frac{1}{16}$ to $\frac{1}{2}$ grain = 0.006 to 0.082 gramme.

It possesses analgesic properties, in 5 p c solutions of the **Sulphate** or **Nitrate** applied locally —*T & G* '85, 376, '86, 18.

Tests —Brucine rapidly loses its Water of crystallisation when exposed to dry air or over Sulphuric Acid at 100° C (212° F) it becomes anhydrous, the anhydrous product melting at 178° C (352.4° F) the aqueous solution is levogyrate, the alkaloid dissolves in concentrated Sulphuric Acid without colour. Concentrated Nitric Acid, or Sulphuric Acid containing Nitric Acid, produces a blood red coloration, passing to orange and finally to yellow. The salts produced when Brucine is neutralised with acid are neutral in reaction towards the customary indicators of neutrality, and the alkaloid may therefore be titrated direct with Normal or Tenth normal Hydrochloric or Sulphuric Acid Solution, using Iodeosin Solution as an indicator of neutrality. 1 c c of Normal Sulphuric or Hydrochloric Acid Solution is equivalent to 0.39133 gramme of anhydrous Brucine or 0.46285 gramme of hydrated Brucine. Brucine should be free from Strychnine, its presence may be detected by oxidising the brucine with Nitric Acid, shaking out the Strychnine by an immiscible solvent and applying the Sulphuric Acid and Potassium Bichromate test when no violet or purple violet coloration should be produced.

OLEA

In the British Pharmacopœia the term **Oleum** is applied to an Oil (whether expressed or distilled), as it is also in Austr., Dutch, Ger., Hung., Jap., Russ. and U.S. The other names for fixed and volatile Oils respectively are Belg., **Oleum** and **Essentia**, Dan.,

Norw and Swed, Oleum and Ætheroleum, Fr, Huile and Essence, Ital Olio and Essenza, Mex, Aceite and Aceite Volatil, Port, Oleo and Essencia, Span, Aceite and Esencia

Elæosaccharum —A title used in the Foreign Pharmacopœias to denote a trituration of an Essential Oil with Sugar Austr, Dutch, Russ and Swiss use 1 drop of the Oil to 2 grammes of Sugar, Belg, Dan and Norw, Oil 1, Sugar 49, Ger, Jap and Swed, Oil 1 gramme, Sugar 50 grammes, they are all practically the same strength Ital (Oleosaccari), Oil 1 gramme, and Sugar 20 grammes, Span, Oil 1, Sugar 25

Not Official

OLEATES

Some of these preparations have come into general use. They were originally made by dissolving the oxide of the metal, or an alkaloid, in an excess of Oleic Acid, but later Dr Shoemaker proposed the method of by double decomposition between a salt of the base and Solution of up (Sodium Oleate with a little Palmitate), Solution of Potassium Oleate may be used with advantage in place of the Solution of Castile Soap, when the pure Oleate is required. The Oleate can also be purified from Palmitate by solution in Petroleum Spirit.

The various Oleates will be found under the headings of their respective bases.

OLIVÆ OLEUM.

OLIVE OIL

FR, HUILE D'OLIVE, GER, OLIVENÖL, ITAL, OLIO DI OLIVE, SPAN, ACEITE DE OLIVAS

A clear pale yellow, or greenish-yellow, oily fluid, possessing a faint characteristic odour and bland oily taste.

It is expressed from the ripe Fruit of *Olea Europæa*.

It is chiefly obtained from the south of Europe.

Adulteration of Olive Oil is very general, large quantities of Cottonseed and other Oils being used for admixture.

On exposure to the air it is apt to become rancid, acquiring a disagreeable smell.

Solubility —1 in 2 of Ether, partially in Alcohol (90 p c)

Medicinal Properties.—Nutritious and mildly laxative, demulcent in the form of emulsion, externally as a lubricant in massage, also as an emollient and protective for burns and certain cutaneous diseases. 4 to 8 fl oz daily, and also larger quantities, have been given in cases of gall stones. Used as a laxative enema, especially for intestinal obstruction (5 oz warm Oil, with or without 8 oz warm Starch Mucilage). Given by the mouth in corrosive poisoning. It is most extensively employed in pharmacy, in the preparation of certain liniments, ointments and plasters.

Its use in typhoid is regarded (*BMJ* '05, 1 414) as a perfect boon. A breakfast cupful is administered as an injection by the bowel once in four or five days at intervals of 12 to 24 hours, and subsequently every second day $\frac{1}{2}$ to 1 fl oz every four hours may also be given by the mouth without producing nausea.

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc, or more

Prescribing Notes—*It may be given as capsules, or in emulsion 1 oz. of Olive Oil with 180 grains of powdered Gum Acacia and Water to 2 oz. Olive Oil mixes well with Malt Extract Heated to 120° to 140° C in a small flask (plugged with Cotton Wool) for half an hour, it forms Oleum Asepticum or Sterilised Olive Oil Almond Oil and Liquid Paraffin can be sterilised in a similar manner*

Official Preparations—Used in the preparation of Emplastrum Ammoniaci cum Hydrargyro, Emplastrum Hydrargyri, Emplastrum Plumbi, Emplastrum Plumbi, Linimentum Ammoniaci, Linimentum Calcis, Linimentum Camphorae, Sapo Durus, Sapo Molli, Unguentum Capsici, Unguentum Hydrargyri Compositum, Unguentum Hydrargyri Nitratis, and Unguentum Resinae

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex (Aceite de Olivo), Norw, Port (Azeite), Russ, Span (Aceite), Swed, Swiss and U.S. Ger and Russ have also Oleum Olivarum Commune Fr has also Huile d'Olive purifiée et stérilisée

Tests—Olive Oil has a sp gr of 0.915 to 0.918. Five samples examined in the author's laboratory had sp gr of 0.916 to 0.917, averaging 0.916. The *B.P.* gravities are 0.914 to 0.919, the *U.S.P.* 0.910 to 0.915 at 25° C (77° F), and the *P.G.* 0.915 to 0.918. It is officially stated to be liable to assume a pasty consistency at 10° C (50° F) and at 0° C (32° F) to form a nearly solid granular mass. The *U.S.P.* states that when cooled from 8° to 10° C (46.4° to 50° F) it becomes somewhat cloudy from separation of crystalline particles, and at 0° C (32° F) it forms a whitish granular mass. The *P.G.* statement is essentially the same as that of the British Pharmacopœia. The congealing point depends greatly upon the length of time to which the Oil is exposed to cold, for instance, the Oil cooled by Ether to -12.8° C (9° F) remained unchanged, but when kept at 0° C (32° F) for 4 hours it partially solidified. Some samples of Oil pressed in the author's laboratory from Olives grown in the south of France showed no sign of congelation during 6 hours at 0° C (32° F) or 3 hours at -9.4° C (15° F). On the other hand, in the following year an Oil from the same district (guaranteed pure) set at once when cooled to -10.6° C (13° F) and within 2 hours at 0° C (32° F). It has since been discovered that the non-freezing Oil is only produced when the fruits have been allowed to over-ripen. The Saponification value and the Iodine absorption afford a useful means of judging the purity of an Olive Oil, but neither are referred to in the *B.P.* The Saponification value should be about 190, the Iodine absorption not less than 80. The *U.S.P.* gives the Saponification value of 191 to 195 and an Iodine absorption of not less than 80 nor more than 88. The *P.G.* makes no reference to the Saponification value, but gives an Iodine absorption of not less than 80 and not more than 84. Five samples of genuine Oil examined in the author's laboratory gave from 189.7 to 198.3 for the Saponification value, with an average of 194.1 and 81.28 to 83.82, with an average of 82.80 for the Iodine absorption. Five other samples of genuine Oil examined for their Iodine value alone showed from 81.28 to 82.82, with an average of 82. Adulteration of Olive Oil is very general, large quantities of Cottonseed and other Seed Oils, Sesame Oil and other Oils being used for the admixture. The *B.P.* includes a test for Cottonseed Oil which is performed by shaking a measured quantity

of 10 c.c. of the Oil with 2 c.c. of a mixture containing 1 p.c. solution of Silver Nitrate in Absolute Alcohol, to which is added 20 p.c. w/v of Ether and a drop of Nitric Acid. It is officially required that no blackening should occur when the mixture is heated on a water-bath for 10 minutes. The *USP* employs an alcoholic solution of Silver Nitrate acidulated with Nitric Acid as described below, but it contains no Ether. The *USP* contains a useful test for the detection of Cottonseed, and is essentially that given in the 17th Edition of the *Companion* and which was suggested to the author by Mr E. J. Bevan and described under Adeps. It consists in heating a few c.c. of the Oil with 1 c.c. of a 1 p.c. solution of Sulphur in Carbon Bisulphide in a salt-bath for about half an hour, no reddish colour should be developed. The *BP* test is essentially Bechi's Silver Nitrate test, it is more generally carried out on the fatty acids of the Oil, and not on the glycerides. The *USP* and *PG* include an Elaidin test with Nitric Acid, which is described in the small type below, this forms a useful means of detecting sophisticated Oil.

The Sugar test is adopted by the *USP* for the detection of Sesame Oil, a 1 p.c. solution of Sugar in Hydrochloric Acid (sp. gr. 1.18) being employed; the test is described below. It frequently contains an excessive amount of free fatty acid, but no test for it is included in *BP*, *USP* or *PG*. The free acid may be determined by warming 5 or 10 grammes of the Oil with 25 c.c. of Alcohol (90 p.c.) cooling and titrating the alcoholic solution with Tenth-normal Volumetric Potassium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 c.c. of Tenth-normal Volumetric Alkali Solution represents 0.028014 gramme of Oleic Acid. The above-mentioned five samples showed from a mere trace to 1.4 p.c. of free acid, with an average of 0.875 p.c. Mineral Oil, if present, may be determined by the amount of unsaponifiable residue, the Oil is saponified with Alcoholic Potassium Hydroxide Solution, evaporated to dryness to remove the Alcohol, the residue is dissolved in Water and the unsaponifiable Oil shaken out with Ether, the ethereal solution evaporated, the residue dried at 100° to 105° C (212° to 221° F) till constant, the residue cooled and weighed. Cottonseed, Rape, or Linseed Oils may be detected by the increased Iodine absorption, as also may Fish Oils. The m.p. of the fatty acid obtained from the Oil also affords a useful indication of the nature of the adulteration. Arachis Oil, which gives an Elaidin test very similar to Olive Oil, may be detected by the isolation of Arachidic Acid.

A test for the absence of Sesame Oil has been suggested with Pyrogallol Solution, 10 c.c. of the Oil are shaken with 10 c.c. of a freshly prepared solution of Pyrogallol (2 grammes) in Hydrochloric Acid (30 grammes) and the separated acid liquid heated in a water-bath for 10 minutes, no distinct violet coloration should be produced.

Nitric Acid—On vigorously shaking 2 c.c. of the Oil and 2 c.c. of Nitric Acid (sp. gr. 1.37), the Oil should retain a light yellow colour, or becoming orange or reddish-brown, and after 6 hours should change into a yellow or white solid mass and an almost colourless liquid, *USP*, 1 c.c. of fuming Nitric

Acid, 1 c.c. of Water and 2 c.c. of Oil at 10° C (50° F), a greenish white, but not red or brown mixture is obtained, which separates into a firm white mass and a faintly coloured liquid after from 2 to 6 hours, *P G*

Silver Nitrate—If 5 c.c. of the Oil be shaken with 5 c.c. of a solution of 0.1 gramme of Silver Nitrate in 10 c.c. of Alcohol, with the addition of 2 drops of Nitric Acid and the mixture heated for about 5 minutes on a water bath, the Oil should retain its original pale colour, not becoming reddish or brown, nor should any dark colour be produced at the line of contact of the two liquids, *U S P*

Amyl Alcohol—If 2 c.c. of the Oil be mixed in a test tube with 2 c.c. of equal volumes of Amyl Alcohol and Carbon Disulphide containing 1 p.c. of Sulphur in solution, and the test tube be immersed to one third or one half its depth in boiling salt Water, no reddish colour should develop in from 10 to 15 minutes, *U S P*

Hydrochloric Acid with Sugar—If a mixture of 2 c.c. of Oil and 1 c.c. of Hydrochloric Acid (sp. gr. 1.18) containing 1 p.c. of Sugar be shaken for half a minute and allowed to stand for 5 minutes, then 3 c.c. of Water added and the whole again shaken, the acid layer should not show a pink colour, *U S P*

OLEUM ARACHIS *Syn.* Earth Nut, Ground Nut or Pea Nut Oil—The oil expressed from the seeds of *Arachis Hypogea*, *Oleum Sesami*, the oil expressed from the seeds *Sesamum indicum*, both are official in the *Ind* and *Col. Adm.*, the former for India and the African, Eastern and Australian Colonies, the latter for India and African, Eastern and North American Colonies, in which places they are officially permitted to be used in place of Olive Oil in making liniments, ointments, plasters and soaps

OPIUM

OPIUM

FR, OPIUM DE SMIRNE, GER, OPIUM, ITAL, OPIO, SPAN, OPTO

The milky exudation of *Papaver somniferum*, L., obtained by incision from the unripe Capsules, and inspissated by spontaneous evaporation

Opium in powder should contain between 9½ and 10½ p.c. of anhydrous Morphine

The Extract and Tincture of Opium being standardised preparations, any suitable variety of Opium may be used in their manufacture, provided that when dry it shall yield when assayed by the official process not less than 7.5 p.c. of anhydrous Morphine. When used in the preparation of the remaining official galenical preparations, Opium is officially required to be of such a strength that the powder obtained from the Opium when dried till constant in weight at 100° C (212° F) shall yield not less than 9.5 and not more than 10.5 p.c. of anhydrous Morphine. The *B P* also permits the dilution of an Opium of greater alkaloidal strength than official requirements to be diluted with one of a less official strength or with Milk Sugar. The *U S P* requires that Opium shall yield when in its normal moist condition not less than 9 p.c. of crystalline Morphine, the *P G* requires that 100 parts of powdered Opium shall contain 10 to 12 parts of anhydrous Morphine, and on drying at 100° C (212° F) shall lose not more than 8 p.c. of its weight.

The Opium official in the *F. Codex* (1908), when dried at 60° C (140° F), is required to contain at least 10 p.c. of Morphine

Opium Granulatum (Opium dried and in coarse powder), and Opium Deodoratum should yield not less than 12 p c nor more than 12.5 p c of crystallised Morphine

Medicinal Properties.—As a hypnotic and sedative it is used in insomnia, excitement and delirium of whatever origin, in the treatment of typhoid, as an analgesic to relieve all forms of neuralgic and abdominal pain, the pain of pleurisy, and of gastric ulcer and of cancer, the pain during the passage of biliary and renal calculi, and the after-pains of labour, as a hæmostatic in intestinal and pulmonary hæmorrhage, in diabetes, in full doses for acute peritonitis, in small doses along with other in diarrhœa

In aortic regurgitation it increases the peripheral blood supply, especially to the brain, it reduces the tendency to syncope, it relieves the angina, and the cardiac dyspnoea, but if the kidneys are affected it should not be given

As an expectorant it is used, guarded by Ammonia, only where the secretion of mucus is abundant, and not thick and viscid or scanty

As a diaphoretic, in form of Dover's Powder, it is valuable in influenza and coryza

As an antispasmodic, in puerperal convulsions, epilepsy, colic, severe forms of chorea and spasmodic asthma, in spasmodic urethral stricture

Locally in the form of liniment, plaster, or fomentation, it is used in neuralgias, rheumatism, and sciatica

To avoid impairment of digestion, and to obtain rapid action, it is given subcutaneously (as hypodermic injection of Morphine) in neuralgia and sciatica, near the seat of pain, also in angina pectoris, cardiac paroxysmal pain, and for the dyspnoea caused by intra-thoracic tumours

In form of Morphine, or Lead and Opium, suppository it relieves rectal and other pelvic pains, and is useful after operations on these regions. Opium is preferable to Morphine in peritonitis, enteritis, and other abdominal inflammations, on account of its direct and more prolonged anodyne and sedative effect, and because of its more continued action it is preferable in delirium and other 'head symptoms'

Its continued use impairs the appetite, digestion and intellect, that it is a cardiac depressant should always be borne in mind. Great caution should be exercised in giving Opium to infants and young children, as they are very susceptible to its action, and it is contraindicated in the pain of chronic dyspepsia, in cases of coma with contracted pupil, in kidney diseases, in nursing females and plethoric persons, in cerebral hyperæmia, in alcoholic intoxication, and for the control of nausea and vomiting in uræmia, in the advanced stages of bronchitis and pneumonia, or whenever the respiration is seriously embarrassed, it is a most dangerous remedy

Valuable papers on Morphine in cardiac diseases—*L* '98, ii 1393, and by Burney Yeo, Stockman, etc on Opium in acute and chronic disease—*Pr* '07, i 626.

Of sugar reducing drugs, the most to be relied upon Most useful in severe cases, in which a rigid diet fails — *Pr* '07, ii 148

A modification of the Biomide treatment of epilepsy is found in the Opium biomide therapy One of the preparations of Opium, preferably the Extract, is given (*L* '05, i 710) for a period of six weeks in increasing doses up to 15 grains per diem, when it is suddenly stopped and large doses of Biomide salt, from 90 to 120 grains, are substituted, this large dose being gradually diminished until about 30 grains are taken daily

A useful way of giving Opium consists (*B M J* '05, ii 1004) in mixing $\frac{1}{2}$ drm to 1 dm of Tincture or Liquid Extract with enough Water to bring it up to 2 fl drm and inject it into the empty rectum by a Glycerin syringe In half to three quarters of an hour the Opium is usually absorbed and relieves pain almost more efficiently and for a longer time than a subcutaneous injection does

Dose — $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Ph Ger maximum single dose, 0.15 gramme, maximum daily dose, 0.5 gramme

Prescribing Notes — *Powdered Opium can be made into pills with Alcohol (60 p c)*

It is convenient to remember that $\frac{1}{16}$ grain Morphine is contained in 1 grain of Powdered Opium, in $\frac{1}{2}$ grain of Extract, in 15 minims of Liquid Extract or of Tincture, in 96 minims of Ammoniated Tincture of Opium, in 240 minims of Compound Tincture of Camphor

Opium is frequently ordered in lotions, 20 to 60 minims of Liquid Extract or Tincture to the fl oz It is also prescribed with Lead Acetate and Lead Subacetate, but the result is a turbid liquid deficient in strength of Lead owing to the precipitation of Lead Meconate, Solution of Morphine Acetate being nearly the same strength as the Tincture, and mixing readily with Lead Lotions without precipitation, can advantageously be employed in its place

Incompatibles — The Alkaline Carbonates, Lime Water, salts of Lead, Iron, Copper, Mercury, and Zinc, Liquor Aisenicalis, and vegetable astringents

Official Preparations — Extractum Opii and Tinctura Opii, used in the preparation of Codeine and of Morphine, of the Powdered Opium, Emplastrum Opii, Pilula Plumbi cum Opio, Pulvis Cretæ Aromaticus cum Opio, Pulvis Opii Compositus, and Unguentum Gallæ cum Opio Contained in Pilula Saponis Composita, Pulvis Kino Compositus, Pulvis Ipecacuanhæ Compositus, and Suppositoria Plumbi Composita Of the Compound Powder, Pilula Ipecacuanhæ cum Scilla, of the Extract, Extractum Opii Liquidum Of the Tincture, Linimentum Opii and Tinctura Opii Ammoniata, contained in Tinctura Camphoræ Composita

Not Official — Acetum, Aqua, Confectio, Enema, Trochiscus, Unguentum, and Vinum, Opii, Solution of Bimeconate of Morphine (Squibb), *Syn* Liquor Meconicus, Meconii Periodidum, Liquor Opii Sedativus, Linctus Opiatus, Linimentum Opii Ammoniatum, Sydenham's Laudanum, Tinctura Opii Clodata, Tinctura Opii Deodorati, Narceina, Narcotina, Papaverina, Cotinine Hydrochloride, Stypticin, and Styptol

Antidotes — In poisoning by Opium the antidotes are, an emetic of 10 grains of Copper Sulphate, the stomach-tube, external stimulants, cold affusion, Ammonia to the nostrils, compelled exertion, and artificial respiration Belladonna or hypodermic injection of Atropine should be used, Strychnine, Amyl Nitrite, Gelsemium, Potassium Permanganate See also Morphine Hydrochloridum

Foreign Pharmacopœias — Official in Austr, not less than 12 p c, Belg, Dan, Dutch, Hung, Ital, Norw, Port and Span, not less than 10 p c, Mex, 10 p c, Ger, Russ, Swed and Swiss, 10 to 12 p c, Fr and Jap, 10 to 11 p c, U S, not less than 12 p c, all calculated on dried Opium

Descriptive Notes — The Opium chiefly imported into this country comes from Asia Minor, Greece, and Persia The Opium from Turkey used in pharmacy is largely that from the districts Karahissar,

Boghaditz and Ghiveh, which latter is often called Constantinople Opium. It occurs in more or less rounded pieces, $\frac{1}{2}$ to 2 lb in weight, usually covered with Poppy leaves, capsules more or less adherent to the cakes. The Boghaditz is richest in Morphine, but gummy and difficult to assay. The Yerli Opium comes from the country surrounding Smyrna, it is soft and unsightly, and chiefly used in the manufacture of Morphine. Natural Karahissar Opium usually yields $11\frac{1}{2}$ to 12 p c of Morphine, it is also used for the manufacture of Morphine, the inferior qualities are known as Adet, Grecian or Salonica Opium, like the two following, is sold as 'shipping' Opium. The best is selected for Cuba, and soft grades are shipped to the United States. It contains from 10 to 14 p c of Morphine, and is sold at a high price for smoking. Tokat and Malatia Opiums are produced in Armenia and go chiefly to Cuba, the West Indies, and Central and South America. They average from 7 to 14 p c of Morphine. These shipping Opiums are generally in softer and flatter cakes and have, on arrival a greener leaf on their surface. When purchased fresh on the Turkey market an allowance is made for the moisture in Opium and a charge of 2d per lb is made for drying it. It loses approximately 23 p c in the drying warehouses before it is fit for shipment. The Boghaditz, Karahissar and Yerli Opiums vary much in size, shape, and weight, and the Poppy leaves covering them are irregularly placed, in the Ghiveh Opium, on the contrary, two leaves are placed in opposite directions on either side of the bun-shaped cakes. All Turkey Opiums have a granular fracture, and consist of agglomerated tears. Persian Opiums, on the contrary, have a uniform non-granular consistence. The Opiums used in pharmacy, although averaging from 11 to 12 p c of Morphine, have of late years, since a uniform standard has become official (10 p c, *BP*), been purposely lowered in Morphine contents. Persian Opium is chiefly re-exported, although it is also used for the manufacture of Morphine, as it averages about 12 p c of that alkaloid, whereas that exported direct from Turkey to China averages 9 to 10 p c, consisting of 80 p c of pure juice and 20 p c of foreign substances. Persian Opium is prepared in various forms, cones, loaves, rectangular blocks, sticks, etc., and these are packed in coloured paper, vine, or fig leaves, sometimes in 'poppy trash,' but the pieces of each brand are usually of uniform size and weight. The Opium in sticks is used for eating, and contains rarely more than 3 p c of Morphine, sometimes only traces. In estimating the value of a chest of Opium, a small portion is taken out of a third of the pieces in the chest, this is beaten into a uniform mass and a small portion of the mass is analysed. The cakes of Opium naturally vary in Morphine contents according to the amount of adulteration with foreign matters and the condition of collection.

Opium is sometimes adulterated with paste made of evaporated grape juice and paste made of dried apricots and inferior gum tragacanth, but such pieces are deficient in elasticity (or 'touch' as the Chinese call it) and break with a short fracture. Particles of Lead added to increase weight have also been found in Opium.

Tests—The *BP* process for the determination of Morphine is a combined gravimetric and volumetric one, the *USP* is a gravimetric process, the Morphine crystals obtained being purified by re-solution in Lime Water and the amount of insoluble matter deducted from the weight of impure Morphine first obtained. The Morphine crystals so obtained contain the Water of crystallisation and are not anhydrous, the *PG* is a volumetric process. The process adopted by the *F¹ Codex* for the determination of the Morphine is the Lime and Ammonium Chloride method, somewhat similar to the *BP*. The crystals are dried at a temperature of 100° C (212° F), and when completely dried (which is stated to require about 2 hours), they are cooled and washed with three successive quantities each of 8 cc of Benzene and again dried at 100° C (212° F), the yield should not be less than 10 nor more than 11 p c. The process at present official in the *BP* is a modification of that of the *BP* 1885, it was originally devised by Portes and Linglois, and with slight alterations was adopted by the Société de Pharmacie de Paris, and was the official process of the *USP* 1880. It was improved by Conroy (*PJ* [3] xv 473) and adopted as the official process in the *BP* 1885. It is a Lime process, and differs only in the following essential points from that of the *USP* 1880: (1) Double the quantities of Opium, Calcium Hydroxide and Ammonium Chloride are employed by the *BP*, and therefore double the quantities are used throughout, (2) the anhydrous and not the crystalline Morphine is weighed, (3) a volumetric determination has been added by the *BP* for the purpose of determining the quantity of pure alkaloid present. The process is almost a verbatim copy of the *USP* 1880, which reads as follows: A weighed quantity of 7 grammes of Opium, in any condition to be valued (the *BP* directs dried at 100° C (212° F) in No. 50 powder), is triturated with 3 grammes of freshly-slaked Lime and 20 cc of Distilled Water in a mortar until a uniform mixture results. A measured quantity of 50 cc of Distilled Water is then added and the mixture stirred occasionally during half an hour, it is filtered through a plated filter into a wide-mouth stoppered bottle having a capacity of about 120 cc, and marked at exactly 50 cc. The *BP* makes an allowance for the soluble matters contained in the Opium, and requires the bottle to be marked at 104 cc, and the corresponding mark on the bottle would therefore be 52 cc. A measured quantity of 50 cc of the filtrate (*BP* quantities correspond to 52 cc), is collected, representing 5 grammes of Opium, 5 cc of Alcohol (94 p c) and 25 cc of Ether added and the mixture shaken, a weighed quantity of 3 grammes of Ammonium Chloride is added and the mixture well and frequently shaken during half an hour, and then set aside for 12 hours to allow the crystallisation of the Morphine. Counterbalance 2 small filters, place one within the other in a small funnel, and decant the ethereal layer as completely as practicable upon the filter. The *BP* inserts instructions as to how the filter papers are to be placed in the funnel, the triple fold of one paper being superimposed upon the single fold of the other. A measured quantity of 10 cc of Ether

is added to the contents of the bottle, which is rotated, and the ethereal layer is again decanted upon the filter, the latter being washed with 5 cc of Ether added slowly and in portions, the filter is now allowed to dry in the air and the liquid in the bottle poured upon it in portions in such a way as to transfer the greater part of the crystals to the filter. The bottle is washed and the remaining crystals are transferred with several small portions of Distilled Water, using not much more than 10 cc in all and distributing the portions evenly upon the filter. The *BP* employs Morphinated Water for washing out the bottles and for washing the crystals on the filter. The Morphinated Water consists of a saturated solution of Morphine in Chloroform Water, and is prepared by digesting an excess of pure Morphine with Chloroform Water for 7 days at a temperature of 15.5°C (60°F). The filter is allowed to drain and dry, first by pressing it between sheets of bibulous paper, afterwards at a temperature between 55° and 60°C (131° to 140°F), this yields the Morphine as a crystalline product containing 1 molecule of Water of crystallisation, the *BP* removes this molecule of Water of crystallisation by further drying for 2 hours at a temperature of 110°C (230°F), and weighs the Morphine in the anhydrous condition, making an allowance of 0.104 of a gramme for each 104 cc for the solubility of Morphine in the menstruum used. Inasmuch as the *BP* process is carried out on twice the quantity recommended by the *U.S.P.* 1880 the weight of anhydrous Morphine obtained plus the solubility allowance of 0.104 of a gramme multiplied by 10 yields the 'cc.' of impure Morphine present in the sample. The *BP* the determination of the amount of pure alkaloid present by the following volumetric method. A weighed quantity of 0.5 gramme of the crystals is titrated with Tenth-normal Volumetric Sulphuric Acid Solution, the titration being officially directed to be continued until the liquid, after boiling, slightly reddens blue Litmus paper. 1 cc of Tenth-normal Volumetric Sulphuric Acid Solution represents 0.0283 gramme of pure anhydrous Morphine. If the number of cc of Tenth-normal Volumetric Sulphuric Acid Solution used be multiplied by 0.0283 the product will be the weight of pure anhydrous Morphine present in the 0.5 gramme of the crystals operated on. From this weight the amount of pure anhydrous Morphine present in the total weight of crystals obtained in the gravimetric process may be calculated, and the resultant weight of pure anhydrous Morphine, plus 0.104 of a gramme (solubility allowance), indicates the amount of pure anhydrous Morphine present in 10 grammes of the Opium, which should amount to not less than 0.95 of a gramme, and not more than 1.05 grammes, corresponding to not less than 9.5 nor more than 10.5 per cent of pure anhydrous Morphine in the dry, powdered Opium. The volumetric portion of the Pharmacopœia process is not very happy. It is better to dissolve a given weight of the crystals in an excess of Tenth-normal Volumetric Sulphuric Acid Solution and to titrate the excess of Tenth-normal Volumetric Acid Solution with Tenth-normal Volumetric Potassium or Sodium Hydroxide Solution, and to use

Methyl Orange Solution in the place of blue Litmus paper as an indicator of neutrality. The part of the test relating to the titration is not very clearly worded, one does not add 'to the weight of anhydrous Morphine indicated by the titration, but to the total weight of the crystals in the filter, afforded by the titration figure'.

The following method of assay is recommended by Dott — A weighed quantity of 10 grammes of powdered Opium is digested with 25 c c of Water, 1.8 grammes of Barium Chloride dissolved in about 12 c c of Water is added, the solution made up to 50 c c, well mixed and after a short time filtered. A measured quantity of 25 c c (= 5 grammes of Opium) is mixed with diluted Sulphuric Acid, just enough to precipitate the Barium, about 1 c c is required, and the solution should be warmed to cause the precipitate to subside, and the solution to filter clear. To the filtered solution about 0.5 c c of dilute Ammonia Solution is added, sufficient to neutralise the free acid, and the solution concentrated to 6 or 7 c c and allowed to cool. 1 c c of Alcohol (90 p c) and 1 c c of Ether is added and Ammonia Solution in slight excess, the Ammonia Solution being added gradually until there is no further precipitation and a perceptible odour of Ammonia remains after well stirring, breaking down any lumps with a stirring-rod. After 3 hours the precipitate is collected on counterpoised filters and washed. It should be noticed that the solution has a faint odour of Ammonia, if not, 1 or 2 drops of Ammonia Solution should be added. The dried precipitate is washed with Benzene or Chloroform, dried and weighed. It is then titrated with Tenth-normal Volumetric Sulphuric Acid Solution until the Morphine is neutralised as indicated by the solution reddening blue Litmus paper. 1 c c of Tenth-normal Sulphuric Acid Solution equals 0.0303 gramme of hydrated Morphine, equivalent to 0.0283 gramme of anhydrous Morphine.

The *USP* process is practically on the following lines — A weighed quantity of 10 grammes of Opium in any condition to be valued is introduced into an Erlenmeyer flask together with 100 c c of Water and the mixture shaken for 10 minutes during 3 hours, the contents are poured on to a wetted filter and when the liquid is drained off, the residue is carefully washed with Water until 150 c c of the filtrate have been obtained, the Water being dropped upon the edges of the filter and its contents. The residual Opium is re-transferred to the flask, 50 c c of Water added, the agitation repeated during 15 minutes and again filtered. The residue is washed as before until a second 150 c c have been collected. The filtrates are evaporated down in rotation in a tared dish, the containing vessels being rinsed out with a third filtrate and the evaporation continued until the residue is reduced to a weight of 14 grammes. After dissolving in the fluid any extract which may have dried on the sides of the basin, it is transferred to a tared Erlenmeyer flask of the capacity of about 100 c c, the dish rinsed with a few drops of Water and the washings transferred to the flask until the mixed solution and washings weigh 20 grammes. 10 grammes of Alcohol (94.9 p c) are added, the flask well shaken and 25 c c of Ether added, the flask

again shaken, 35 c.c. of Ammonia Solution are now added from a graduated pipette or burette, the flask stoppered and shaken thoroughly during 10 minutes and set aside in a moderately cool place for at least sixteen hours. The ethereal solution is decanted as completely as possible on to two small counterpoised filters contained in a glass funnel in such a way that the triple fold of the inner filter is laid against the single fold of the outer filter, both being previously moistened with Ether. The contents of the flask are washed with 10 c.c. of Ether, which is also decanted on to the filter, the operation being completed with a further quantity of 10 c.c. of Ether, the filter paper is then dried, and the aqueous contents of the flask are then transferred to the filter, the crystals of Morphine, and the aqueous contents of the bottle are transferred to the filter, the remaining crystals being removed from the flask with Water, using not more than 15 c.c. in all. The filter is allowed to drain, washed with Alcohol (94.9 p.c.) previously saturated with powdered Morphine, and finally with Ether, using about 10 c.c. or more if necessary. The filter is allowed to dry at a temperature not exceeding 60° C (140° F) until its weight remains constant, transferred to a tared watch-glass and weighed. The crystals are placed in an Erlenmeyer flask together with 10 c.c. of Lime Water for each 0.1 of a gramme of Morphine and the mixture shaken at intervals during 30 minutes. The liquid is passed through two filter papers folded so that the triple fold of the inner filter paper is superimposed against the outer filter paper, the flask rinsed with Lime Water and the washings passed through the filter until the filtrate, after acidification, no longer yields a precipitate with Mercuric Potassium Iodide (Mayer's) Solution, the filters are pressed between folds of bibulous paper until nearly dry, and dried to a constant weight. The weight of the insoluble matter on the filter deducted from the weight of Morphine crystals previously found, and the difference multiplied by 10, represents the percentage of crystallised Morphine contained in the Opium.

The *PG* method of determination is as follows.—A weighed quantity of 6 grammes of Opium in a state of middling fine powder is triturated with 6 grammes of Water and the mixture transferred to a dry tared flask and the contents brought to a weight of 54 grammes by the addition of a further quantity of Water. After the mixture has been allowed to stand for 1 hour with intervals of vigorous shaking, the mass is pressed through a piece of dry calico, 42 grammes of the pressed fluid filtered through a dry filter paper into a dry flask, 2 grammes of a 1 in 2 w/w Sodium Salicylate Solution added and the whole vigorously shaken. A weighed quantity of 36 of the clear fluid is then filtered through a dry filter into a flask, the filtrate is rotated with 10 grammes of Ether and mixed with 5 grammes of a mixture of 17 grammes of Ammonia Solution and 83 grammes of Water, the flask is then closely stoppered, the contents shaken vigorously for 10 minutes and allowed to remain at least for 24 hours. The ethereal liquid is then completely transferred to a plated filter, the aqueous liquid remaining in the flask is washed with 10 grammes of Ether, the mixture allowed to remain a few seconds and then trans-

ferred again to the filter. After the separation of the ethereal liquid the aqueous solution is transferred through the same filter together with the crystalline residue. The filter as well as the flask is washed 3 times with successive quantities of 5 grammes of Water saturated with Ether, after the flask has been thoroughly washed out and the filter has been completely drained, the Morphine crystals, after drying, are dissolved in 25 cc of Tenth-normal Volumetric Hydrochloric Acid, the solution is transferred to a flask of 100 cc capacity, the filter and flask washed with Water and the solution diluted to 100 cc. A measured quantity of 50 cc of this solution is transferred to a stoppered flask of about 200 cc capacity, 50 cc of Water added, and sufficient Ether added to form a layer of about 1 cm, and the excess of Tenth-normal Volumetric Hydrochloric Acid Solution is titrated with Tenth normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodopsisin Solution as an indicator of neutrality, shaking the solution after each addition. Not more than 5.4 cc and not less than 4.1 cc of the Tenth-normal Volumetric Potassium Hydroxide Solution should be required. The number of cc of Tenth-normal Volumetric Potassium Hydroxide Solution required multiplied by 2, the product subtracted from 25, the difference multiplied by 0.0285 yields the weight of anhydrous Morphine present in 4 grammes of the Opium, and if this figure be again multiplied by 25 will yield the percentage by weight of anhydrous Morphine present in the sample.

Preparations

EMPLASTRUM OPII OPIUM PLASTER

Opium in very fine powder, 1, Resin Plaster, 9 (1 in 10)

Anodyne to relieve local pain

Foreign Pharmacopœias—Official in Mex, 1 Opium in 20 Fr, 1 Extract in 4, Port, 1 Extract in 10, U S, 6 Extract in 100. Not in the others.

EXTRACTUM OPII. EXTRACT OF OPIUM

An Aqueous Extract containing 20 p c of Morphine

Dose— $\frac{1}{4}$ to 1 grain = 0.016 to 0.065 gramme

Ph Ger maximum single dose, 0.15 gramme, maximum daily dose, 0.5 gramme

The *B P* and *U S P* both require the Extract to contain 20 p c of Morphine, the *P G* Extract is required to yield from 15 to 20 p c of Morphine.

The Extract of Opium official in the *H^o Codex* (1908) is required to contain exactly 20 p c of Morphine, which is in accordance with the recommendation of the *Brussels Conference*.

The *Brussels Conference* agreed to a content of 20 p c of Morphine in the extract.

It is officially permitted to mix stronger and weaker Extracts in order to obtain extract of Opium of proper strength and consistence. An Extract of Opium stronger than the official requirements may be diluted with a sufficiency of Distilled Water or with Milk Sugar. In the first issue of the *B P* '85 the Extract was directed to be made from Opium in powder and restricted in the official variety, but the criticism evoked was so strong that in the later reprints it was permitted to use any variety of Opium as long as the product conformed to the official standard of Morphine.

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Ger,
2 1 2

Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S Not in Dan

The International Standard is 20 p c of Morphine Austr, Belg, Dutch, Fr, Mex, Span, Swiss and U S all 20 p c of Morphine Ger, 15 to 20 p c Jap, 16 to 17 5 p c Ital, 15 p c Swed, 17 to 20 p c The remainder do not give percentage

Tests—Extract of Opium is assayed by the process described under Opium, a weighed quantity of 7 grammes of the Extract being used in place of the 14 grammes of powdered Opium there employed. The *USP* employs the following process for the assay of the Extract—A weighed quantity of 4 grammes is dissolved in 30 c c of Water, the solution filtered, the filter and residue washed with Water until all soluble matter is extracted, the washings being separately collected. They are evaporated in a tared dish to a weight of 10 grammes. When completely dissolved the Extract is poured into a tared Erlenmeyer flask of about 100 c c capacity, the dish rinsed with a few drops of Water until the total weight of the solution amounts to 15 grammes, a weighed quantity of 7 grammes of Alcohol (94.9 p c) is added, the flask well shaken and 20 c c of Ether added, the shaking being repeated. 22 c c of Ammonia Water is added from a pipette, the flask stoppered and thoroughly shaken for 10 minutes and set aside for 6 hours or over night. The Ether solution is then filtered through two counterpoised filter papers, the filter papers being placed in the funnel in such a manner that the triple fold of the inner filter is superposed on the single fold of the outer filter, and the papers are previously moistened with Ether. The contents of the flask are washed with 15 c c of Ether, which Ether solution is again decanted on the filter, the washing is repeated with a further portion of 15 c c of Ether, the aqueous liquids in the flask are transferred to the filter together with the crystals of Morphine, the crystals remaining in the flask being transferred by washing with several portions of Water, using a total quantity of not more than 10 c c. The filter is allowed to drain, the crystals being washed free from the mother liquor first with Water and afterwards with (94.9 p c) Alcohol (saturated with Morphine), and finally with Ether, using about 10 c c or more if necessary. The filter is allowed to dry at a temperature not exceeding 60° C (140° F) until constant in weight, the crystals of Morphine carefully transferred to a watch-glass and weighed, they are then transferred to an Erlenmeyer flask, and Lime Water in the proportion of 10 c c for each 0.1 of a gramme of Morphine added, the flask being shaken at intervals for 25 minutes, the filter placed in a funnel in such a way that the triple fold of the inner filter is laid against the single fold of the outer filter, the flask is rinsed with Lime Water passing the washings through the filter until the filtrate after being acidified will no longer yield a precipitate with Mercuric Potassium Iodide (Mayer's) Solution. The filters are pressed between folds of bibulous paper, dried to a constant weight and weighed. The weight of insoluble matter on the filter is subtracted from the weight of the impure Morphine crystals found above, the difference multiplied by 5 yields the percentage of pure crystalline Morphine present in the Extract of Opium.

The German Pharmacopœia employs the following method for the determination of Morphine in the Extract —A weighed quantity of 3 grammes is dissolved in 40 grammes of Water, 2 grammes of a 1 in 2 w/w solution of Sodium Salicylate added to the solution and 30 grammes of the clear liquid filtered through a dry filter into a dry flask. The filtrate is mixed with 10 grammes of Ether and 5 grammes of a mixture of 17 grammes of Ammonia Solution and 83 grammes of Water, the flask is stoppered, the contents shaken vigorously for 10 minutes and allowed to remain at rest for 24 hours. The ethereal liquid is then filtered through a counterpoised filter, the aqueous fluid remaining in the flask is washed with 10 grammes of Ether and the ethereal liquid passed through the same filter. The aqueous solution is then passed through the filter without removing the residue which is crystallised on the sides of the flask, the flask is washed with 3 successive quantities of Water saturated with Ether. After it has been well washed and the filter is completely drained, the Morphine crystals, after drying, are dissolved in 25 cc of Tenth normal Volumetric Hydrochloric Acid, the solution transferred to a flask of 100 cc capacity, the filter and the flask washed thoroughly with Water, and the filtrate and washings finally diluted to 100 cc. A measured quantity of 50 cc of this solution is mixed in a stoppered flask of about 200 cc capacity with 50 cc of Water and sufficient Ether to form a layer of about 1 cm. The excess of acid is titrated with Tenth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosin Solution as an indicator of neutrality. Not more than 6.5 cc and not less than 5.5 cc of the tenth-normal volumetric alkali solution should be necessary to neutralise the excess of acid. The number of cc of Tenth-normal Volumetric Potassium Hydroxide Solution required is multiplied by 2, the product subtracted from 25, the difference is multiplied by 0.0285, and this product multiplied by 100 and divided by 2.25 yields the percentage by weight of anhydrous Morphine in the extract.

EXTRACTUM OPII LIQUIDUM LIQUID EXTRACT OF OPIUM
Extract of Opium, $\frac{3}{4}$, Distilled Water, 16, Alcohol (90 p c), 4

Contains 1 grain of Extract = $\frac{1}{2}$ grain Morphine, in 29 minims

Dose —5 to 30 minims = 0.3 to 1.8 cc

Not in the foreign Pharmacopœias

The *B P* Liquid Extract of Opium is required to contain 0.75 p c w/v of anhydrous Morphine, a Fluid Extract is not official in either the *U S P* or *P G*

Tests —Liquid Extract of Opium has a sp gr of 0.985 to 0.990, it contains about 3 p c w/v of total solids, and about 18 p c w/v of Absolute Alcohol. It is officially required to contain not less than 0.7 p c w/v nor more than 0.8 p c w/v of anhydrous Morphine as determined by a similar process to that adopted for the determination of the alkaloid in the Tincture (*see Tinctura Opi*)

LINIMENTUM OPII. LINIMENT OF OPIUM

Tincture of Opium, 1, Liniment of Soap, 1 (1 in 2)

The addition of the Opium to the Soap Liniment renders it more useful in many cases of rheumatism and local pains

Official in Span, *Tintura Alcoholica de Opio Jabonosa*

PILULA SAPONIS COMPOSITA. COMPOUND PILL OF SOAP

Opium, in powder, 1, Hard Soap, in powder, 3, Syrup of Glucose (by weight), 1 (1 of Powdered Opium in 5)

Dose—2 to 4 grains = 0.13 to 0.26 gramme

Foreign Pharmacopœias—Official in Fr, 1 Extract in 10, Dan (*Pilulas Cynoglossi*), about 1 in 7, Norw, 1 Opium in 7½, Span, 1 Extract in 10 Part (*Pilulas de Opio Comp*), 1 Extract in 10, US (*Pilula Opium Powdered Opium* 6½, Soap 2, Mex has *Pildoras pacificas*, each containing 0.2 gramme of Opium with other ingredients Not in the others

PULVIS OPII COMPOSITUS COMPOUND POWDER OF OPIUM

Opium, 3, Black Pepper, 4, Ginger, 10, Calaway Fruit, 12, Tiagacanth, 1 (1 of Powdered Opium in 10)

1 of this powder with 3 of Syrup forms *Confectio Opii*, *BP '85*

Dose.—2 to 10 grains = 0.13 to 0.65 gramme

TINCTURA OPII TINCTURE OF OPIUM *BP Syn*—LAUDANUM

N O.Syn—TINCTURA THEBAICA

Opium treated with equal volumes of Distilled Water and Alcohol (90 p.c.), and standardised to contain 0.75 gramme of anhydrous Morphine in 100 c.c.

Contains ½ grain Morphine in 29 minims

Dose—5 to 15 minims = 0.3 to 0.9 c.c., for repeated administration, for a single administration, 20 to 30 minims = 1.2 to 1.8 c.c.

Ph Ger maximum single dose, 1.5 gramme, maximum daily dose, 5.0 grammes

The *BP* Tincture of Opium is required to yield 0.75 p.c. w/v of anhydrous Morphine. The *USP* Tincture is required to contain not less than 1.2 p.c. w/v nor more than 1.25 p.c. w/v of crystallisable Morphine. The Tincture of the German Pharmacopœia is required to yield 1 to 1.2 p.c. w/v of anhydrous Morphine.

The *Brussels Conference* fixed the strength of the Tincture at 10 p.c. of Opium, and Alcohol (70 p.c.) as a menstruum for the preparation of the Tincture, that it shall be prepared by percolation, and that the strength in Morphine should be 1 p.c. w/w. The Tincture of Opium official in the *Fr Codex* conforms to these requirements, but it is made from the standardised Extract.

This preparation is stated officially to contain the soluble matter of 32.8 grains of Opium (containing 10 p.c. of anhydrous Morphine) in 1 fl. oz. or about 1 grain of such Opium in 15 minims.

Provided that the Opium does not contain less than 7.5 p.c. of Morphine calculated as anhydrous, any variety is officially allowed for

the preparation of the Tincture, it being also stipulated that the resultant tincture should correspond to the quantitative test given above

B P '85 ordered a definite quantity of Opium containing about 10 p c of Morphine, only about three quarters of the Morphine was extracted from the Tincture, but the figure for Morphine was fixed on a different assumption. This difficulty is now removed by fixing a standard for the Morphine content of the Tincture irrespective of the quantity of Opium employed

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Ger, Hung, Ital, Jap, Norw, Russ, Swed, Swiss and U S, 1 (powder) in 10, Mex, 1 in 8, Fr, Port and Span, 1 Extract in 20. All by weight, except U S. U S has also *Tinctura Opii Deodorati*

The *Brussels Conference* agreed to a strength of 10 p c of Opium, 1 p c of Morphine, using Alcohol (70 p c)

The *Fr Codex* contains 5 p c w/w of Extract of Opium corresponding to about 10 p c of Opium dried at 60° C (140° F), it is prepared with 70 p c Alcohol in accordance with the recommendation of the *Brussels Conference*

Tests—Tincture of Opium has a sp gr of 0.950 to 0.960, it contains about 3.5 p c w/v of total solids and about 43 to 44 p c w/v of Absolute Alcohol. It is officially required to yield not less than 0.7 p c w/v nor more than 0.8 p c w/v of anhydrous Morphine as determined by evaporating a measured quantity of 80 c c of the Tincture to about $\frac{1}{3}$ th its volume, adding 3 grammes of freshly slaked Lime, and thoroughly mixing and diluting the mixture with Water to 85 c c. It is then set aside, with intervals of occasional shaking, for 30 minutes. A measured quantity of 50 c c (= 50 c c Tincture) is filtered into a wide-mouthed, stoppered bottle, 5 c c of Alcohol (90 p c) and 30 c c of Ether added, and the mixture shaken, 2 grammes of Ammonium Chloride is then added, and the mixture frequently and vigorously shaken during 30 minutes, and finally set aside for 12 hours to allow the Morphine to crystallise. The ethereal liquid is then suitably transferred to two small counterpoised filter papers contained in a funnel in such a way that the triple fold of the one filter shall be laid upon the single fold of the other filter paper. The transference is preferably not made in a pipette as recommended in the *B P*, which is clumsy and apt to result in loss of alkaloid. The aqueous liquid in the bottle is washed by shaking with 15 c c of Ether, which ethereal solution is passed through the same filter paper, and the filter is finally washed with 10 c c of Ether. After the filter has been allowed to dry, the aqueous liquid is filtered through the same filter paper, the crystals being transferred to the filter, first by means of small successive quantities of the filtrate, and the last traces of crystals are transferred from the bottle by washing with Morphinated Water, the crystals on the filter are washed with Morphinated Water until the washings are colourless, dried first by pressure between folds of bibulous paper, subsequently at a temperature not exceeding 60° C (140° F), and finally are rendered anhydrous by drying at 110° C for 2 hours, cooled and weighed. The solubility allowance recommended by the

BP is of 0.05 of a gramme or 0.1 of a gramme for every 100 c.c. of the original filtrate, and this figure must be added to the weight of crystals obtained in the gravimetric determination, the product multiplied by 2, indicates the percentage w/v of anhydrous Morphine present in the Tincture. A weighed quantity of 0.3 gramme of the crystals is titrated with Tenth-normal Volumetric Sulphuric Acid Solution in the same manner as directed for the crystals obtained in the Opium determination. The number of c.c. Tenth-normal Volumetric Sulphuric Acid Solution required, multiplied by 0.0283, represents the amount of pure anhydrous Morphine present in 0.3 of a gramme of the crystals worked upon, from this the weight of pure anhydrous Morphine present in the total amount of crystals obtained in the determination may be calculated, to the weight of pure anhydrous Morphine thus obtained, is added 0.05 of a gramme or 0.1 of a gramme for every 100 c.c. of the original filtrate and the product multiplied by 2 yields the percentage w/v of pure anhydrous Morphine present in the Tincture.

The *USP* method of assay is to evaporate 100 c.c. of the Tincture to about one-fifth of its volume, add 40 c.c. of Water and mix thoroughly, set aside the mixture for 1 hour, stirring occasionally during the interval to disintegrate the resinous flakes adhering to the dish. It is then filtered, the residue washed with Water until all the soluble matter is extracted, and the filtrate and washings evaporated in a tared dish to a weight of 14 grammes, which is then assayed according to the process described under Opium. In calculating the results the final multiplication by 10 is omitted, as the 14 grammes worked upon represents 100 c.c. of the Tincture.

The *PG* evaporates 50 grammes in a weighed porcelain dish to 15 grammes, dilutes with Water to a weight of 38 grammes, adds 2 grammes of a 1 in 2 w/w solution of Sodium Salicylate, and after vigorous shaking filters 32 grammes of the clear fluid through a dry filter paper into a dry flask. The filtrate is shaken with 10 grammes of Ether, 5 grammes of a mixture of 17 grammes of Ammonia Solution and 83 grammes of Water added, the flask is stoppered and the contents vigorously shaken for 10 minutes and allowed to remain at rest for 24 hours. The ethereal layer is then completely separated, passed through a counterpoised filter, the aqueous solution remaining in the flask washed with 10 grammes of Ether, and this ethereal liquid again passed through the filter. After the complete separation of the ethereal fluid, the aqueous solution is passed through the same filter without disturbing the crystalline residue which is attached to the sides of the flask. The filter and flask are washed with 3 successive quantities each of 5 grammes of Water saturated with Ether, and after the flask has been well washed the filter is completely drained. The Morphine crystals, after drying, are dissolved in 25 c.c. of Tenth-normal Volumetric Hydrochloric Acid, the solution transferred to a flask of 100 c.c. capacity. The filter and flask washed with Water and the solution diluted to 100 c.c., a measured quantity of 50 c.c. of this solution is transferred to a stoppered flask of about 200 c.c. capacity, 50 c.c. of Water

added and sufficient Ether to form a layer of about 1 cm. The excess of Tenth-normal Volumetric Sulphuric Acid Solution is titrated with Tenth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosin Solution as an indicator of neutrality. After each addition the mixture is vigorously shaken, not more than 5.5 cc and not less than 4.2 cc of Tenth-normal Volumetric Alkali Solution shall be required. The number of cc of Tenth-normal Volumetric Alkali Solution required should be multiplied by 2, the product subtracted from 25, the difference multiplied by 0.0285, and the product again multiplied by 100 and divided by 40 yields the percentage w/v of anhydrous Morphine present in the Tincture.

TINCTURA OPII AMMONIATA AMMONIATED TINCTURE OF OPIUM Scotch Paregoric

Tincture of Opium, 3 fl oz, Benzoic Acid, 180 grains, Oil of Anise, 1 fl dr, Solution of Ammonia, 4 fl oz, Alcohol (90 pc), *q s* to yield 20 fl oz.

Tincture of Opium is now used instead of Powdered Opium, Saffron is omitted, and Liquor Ammoniae Fortis replaced by Liquor Ammoniae.

Dose — $\frac{1}{2}$ to 1 fl dr = 1.8 to 3.6 cc

Contains $\frac{1}{16}$ grain Morphine in 32 minims

TINCTURA OPII BENZOICA — See TINCTURA CAMPHORAE COMPOSITA

Other preparations containing Opium —

	Proportion of Opium
Pilula Ipecacuanhæ cum Scilla	about 1 in 20
Pilula Plumbi cum Opio	1 in 8
Pulvis Cretæ Aromaticus cum Opio	1 in 40
Pulvis Ipecacuanhæ Compositus	1 in 10
Pulvis Kino Compositus	1 in 20
Suppositoria Plumbi Composita	1 grain in each
Tinctura Camphoræ Composita	$\frac{1}{2}$ grain in 1 fl dr
Unguentum Gallæ cum Opio	1 in 13 $\frac{1}{2}$

	Proportion of Morphine salt
Injectio Morphine Hypodermica	1 in 20
Liquor Morphine Acetatis	1 in 100
Liquor Morphine Hydrochloridi	1 in 100
Liquor Morphine Tartratis	1 in 100
Suppositoria Morphine	$\frac{1}{2}$ grain in each
Tinctura Chloroformi et Morphine Composita	1 in 100
Trochiscus Morphine	$\frac{1}{10}$ grain in each
Trochiscus Morphine et Ipecacuanhæ	$\frac{1}{30}$ grain in each

Not Official

AQUA OPII — Opium, in powder, 1, Water, 12, distil 6
Occasionally employed in eye lotions Aqua Opi, 1, Aqua Sambuci, 7

ACETUM OPII — Powdered Opium, 10, Myristica, in No. 30 powder, 3, Sugar, 20, Diluted Acetic Acid, *q s* to make 100 — *U S P*

Average Dose. — 8 minims = 0.5 cc

Acetum Opi, *B P C*, nearly corresponds to the above, the difference being in the Diluted Acetic Acid, which is stronger in the *U S P* than in the *B P*

CONFECTIO OPII —Compound Powder of Opium, 1, Syrup (by weight), 3
—*B P* 1885

This has been incorporated in the *B P C*

ENEMA OPII —Tincture of Opium, 10 to 40 minims, Mucilage of Starch, 2 to 4 fl oz —*St Thomas's*

Tincture of Opium, 3, Mucilage of Starch, *q s* to make 100 —*B P C*

The quantity sufficient for one application is 2 fl oz, which should be administered warm —*B P C*

LINCTUS OPIATUS —Tincture of Opium, 2 minims, Oxymel of Squill, 15 minims, Mucilage of Tragacanth, 15 minims, Glycerin, 15 minims, Emulsion of Chloroform, 3 minims, Syrup, to 1 fl drn —*St Thomas's*

This has been incorporated in the *B P C*

LINIMENTUM OPII AMMONIATUM —Liniment of Soap, 6, Compound Camphor Liniment, 6, Tincture of Opium, 6, Liniment of Belladonna, 1 Stronger Solution of Ammonia, 1, mix, and after standing a week, filter quickly —*B P C Formulary* 1901, now incorporated in the *B P C*

LIQUOR MORPHINÆ BIMECONATIS (*Squire*) *Syn* **Liquor Meconicus** —A purified Solution of Opium (introduced by Peter Squire in 1839), containing the whole of the alkaloids in their natural state of combination. It is now standardised to contain 1 p c of Morphine. The volatile and extractive matters, to which the unpleasant secondary effects of Opium have been attributed, are removed in the process of its manufacture.

The Solution of the same name inserted in the *B P* of 1885, though obviously intended to take its place, differed so widely from the original in its properties and method of preparation, that it was no substitute for it, and was deleted in 1898.

Dose —5 to 30 minims = 0.3 to 1.8 c c

LIQUOR OPII SEDATIVUS (*Battley*) has enjoyed a reputation for a long time as an anodyne and sedative superior to Tincture of Opium, but it is somewhat stronger.

Dose —5 to 20 minims = 0.3 to 1.2 c c

Liquor Opii Sedativus —Opium (10 p c), 2 oz, Calcium Hydrate, 2 drn, Alcohol (90 p c), 4 oz, Sherry, 3 oz, Water, *q s*. Boil the Opium (broken into small pieces) and Lime in 15 oz of Water for half an hour, and allow to cool. Make up to 13 oz with Water, add the Alcohol and Sherry. Filter, press the marc, add the expressed liquid filtered, and to this add Proof Spirit to make 20 fl oz. Set aside for 6 months to mature, then improved. —*A Ph F*

Opium, in small pieces, 10, Calcium Hydroxide, 1.50, Alcohol, 20, Sherry, 15. Distilled Water, *q s*, Alcohol (60 p c), *q s* to produce 100 —*B P C*.

MECONII PERIODIDUM —A preparation representing the alkaloids of the above preparation in combination with excess of Iodine, on the lines of the other Di-iodo-hydriodides.

Dose — $\frac{1}{8}$ to $\frac{1}{2}$ gram = 0.008 to 0.032 grammes

TINCTURA OPII CROCATI (Sydenham's Laudanum) —Contains Saffron, and occurs in the majority of the foreign Pharmacopœias. The *Brussels Conference* agreed to a strength of 1 p c w/w of Morphine.

All the preparations are by weight, except *B P C*

Austr —Opium 10, Saffron 2, Alcohol (68 p c) 40, Cinnamon Water 60

Dutch —Opium 60, Saffron 20, Cinnamon 5, Cloves 5, Alcohol 250, Water 250.

Macerate for 8 days, express, filter, and, if necessary, dilute to contain 1 p c. of Morphine.

Hung —Opium 15, Saffron 15, Cinnamon Water 150

Ger —Opium 15, Saffron 5, Cloves 1, Cassia 1, Alcohol (70 p c) 70, Water 70.

Russ —Opium 15, Saffron 5, Cloves 1, Cassia 1, Alcohol (70 p c) 70, Water 75.

Swiss —Opium 10, Saffron 5, Cloves 1, Cassia 1, Alcohol (68 p c) 94

Opium 5, Cinnamon Bark 1, Cloves 1, Saffron 5, Detannated Sherry, *q s* to produce 100 — *B P C*

Laudanum Sydenhami

Belg — Extract of Opium 50, Tincture of Saffron 150, Oil of Cinnamon 1, Eugenol 1, Alcohol (70 p c) 798

Laudano de Sydenham

Mex — Opium 10, Saffron 5, Oil of Cinnamon 16 drops, Oil of Cloves 16 drops, Crystallisable Acetic Acid 0.8, Alcohol (30 p c) 80

Laudanum de Sydenham

Fr — Opium 100, Saffron 50, Oil of Cloves 1, Oil of Cinnamon 1, Alcohol 30 p c 1000

It is required to 1 p c of Morphine in conformity with the recommendation of the *Brussels Conference*.

Vinum Opii

U S — Opium 10, Cassia 1, Cloves 1, Alcohol 15, White Wine to measure 100

Vinum Opii Aromaticum

Jap — Saffron 1, Cloves 1, Cinnamon 1, Dilute Spirit 7, Sherry 85, Opium 1

Vinum Opii Crocatum

Norw — Opium 15, Saffron 5, Cloves 1, Cinnamon 1, Malaga Wine 150

Swed — Opium 15, Saffron 5, Cloves 1, Cinnamon 1, Musika Wine 150

Vinho de Opio Composto

Port — Extract of Opium 5, Saffron 3, Cloves 1, Cinnamon 1, Madeira Wine 100

Vino de Opio Compuesto

Span — Opium 10, Saffron 5, Cloves 1, Cinnamon 1, White Wine to 100

Laudano Vini Opiiato Composto

Ital — Opium 16, Saffron 8, Cinnamon 1, Cloves 1, Alcohol (60 p c) 70, Water 70

Tinctura Thebaicum Crocatum

Dan — Opium 100, Saffron 25, Cloves 6, Cinnamon 6, Alcohol (68 p c) to 1000

TINCTURA OPII DEODORATI — Granulated Opium (containing 12 to 12.5 of Crystallisable Morphine), 10, Purified Petroleum Benzine, 7.5, Alcohol (95 p c), 20, Water, *q s* to produce 100. Heat 50 of Water to boiling and pour it on the Granulated Opium contained in a suitable vessel, stirring the mixture frequently during 24 hours. Transfer to a percolator, return the first portion of the percolate until it runs through clear, and when the liquid ceases to drop continue the percolation with Water until the Opium is exhausted. Concentrate the percolate on a water bath to 15, and when cool shake it vigorously for 10 minutes with 6.5 of the Purified Petroleum Benzine, separate the Benzine, repeat the shaking out for a few minutes with the remainder of the Benzine and, having carefully and completely separated this second portion of Benzine, evaporate the remaining liquid in a warm place spontaneously until the odour of Benzine has disappeared, removing the last traces by the heat of a water bath. Mix the deodoured liquid so obtained with 60 of Water, filter the mixture through a paper filter and, having mixed the Alcohol with the filtrate, wash the filter with sufficient Water to make 100 — *U S P*

Average Dose — 8 minims = 0.5 c c

It should contain 1.2 to 1.25 p c w/v of crystallised Morphine

This has been incorporated in the *B P C* in a modified form, using Opium, *B P*, 7.5, adjusting the strength of the Tincture to 0.75 p c of anhydrous Morphine

TROCHISCUS OPII — $\frac{1}{10}$ grain of Extract of Opium in each

Dose — 1 to 6 lozenges

U S, Powdered Opium $\frac{1}{10}$ grain in each

UNGUENTUM OPII — Extract of Opium, 1, Spermaceti Ointment, 9

Rub the Extract with a small quantity of Water to a syrupy consistence, and mix with the Ointment, (1 in 10)

VINUM OPII (sine Aromat)—Opium, in powder, 1, Sherry, 10 Mace-
rate 7 days, and filter

Used as a **collyrium**, 1 to 16 of Water

Dose—10 to 40 minims = 0.6 to 2.4 cc

VINUM OPII—Extract of Opium, 1 oz., Cinnamon Bark, 75 grains,
Cloves, 75 grains, Sherry, 20 fl oz. **Dose**—10 to 40 minims = 0.6 to 2.36 cc
Each fl dram contains about half a grain of Morphine.—*B P* 1885

This has been incorporated in the *B P C*, using detannated Sherry

NARCEINA Narceine $C_{23}H_{27}NO_4$, $3H_2O$, eq 495.55—In white, silky,
acicular crystals, neutral, with a slightly bitter taste. Soluble in 375 parts of
cold and in 220 of hot Water, also soluble in Alcohol, insoluble in Ether, and
practically insoluble in Chloroform

It should be kept in well-stoppered glass bottles of a dark amber tint and
as possible from contact with air, as it is liable to absorb both
and moisture

Dose— $\frac{1}{2}$ to 1 grain = 0.032 to 0.065 gramme

Tests—Narceine should not melt under $165^{\circ}C$ ($329^{\circ}F$). Commercially,
pure Narceine should not fuse under $170^{\circ}C$ ($328^{\circ}F$). It contains three
molecules of Water of crystallisation, equivalent to 10.8 p.c., which are lost at a
temperature of $100^{\circ}C$ ($212^{\circ}F$), and when heated to a still higher temperature
it evolves an odour resembling Trimethylamine. The aqueous solution should
be neutral in reaction towards Litmus paper, the alkaloid dissolves completely
in diluted Sulphuric Acid, and if this acid solution be concentrated on a water-
bath a beautiful violet coloration is produced, which changes to cherry-red. On
further heating upon the introduction of a trace of Nitric Acid, a bluish-violet
streak is produced. It is precipitated by the usual alkaloidal reagents, *e.g.*,
Mercuric Potassium Iodide (Mayer's) Solution, Iodo-Potassium Iodide (Wagner's)
Solution and Ferric Acid Solution. Diluted Iodine Solution
coloration. Narceine may be distinguished from Morphine
yielding a blue coloration with Potassium Ferrocyanide containing a trace of
Ferric Chloride T.S. It should leave no weighable residue when ignited with free
access of air

Foreign Pharmacopœias—Official in Mex. Not in the others

Under the title **Antispasmin**, a combination of Narceine-Sodium and
Sodium Salicylate has been introduced as a hypnotic and sedative

Dose— $\frac{1}{4}$ to 2 grains = 0.016 to 0.13 gramme

Narceyl—In chemical constitution it is an Ethyl-narceine Hydrochloride,
and forms fine silky needles, sparingly soluble in Water. It is stated to be useful
in allaying the severe cough in cases of pulmonary tuberculosis

Dose—1 to $1\frac{1}{2}$ grain = 0.065 to 0.1 gramme, in 24 hours

NARCOTINA Narcotine $C_{22}H_{23}NO_7$, eq 410.12—Trimetric prismatic
crystals, or in large, colourless, glistening needles. Insoluble in Water, soluble
in Ether, in boiling Alcohol, and in diluted Acids, insoluble in Potassium
Hydroxide Solution. It has no narcotic properties, and has therefore been called
Anarcotina, it has been given as a substitute for Quinine, as an antiperiodic
in ague

Dose—1 to 3 grains = 0.06 to 0.2 gramme

Tests—Narcotine melts at about $170^{\circ}C$ ($338^{\circ}F$), and when heated to a
somewhat higher temperature evolves an odour somewhat resembling Trimethyl-
amine. It dissolves completely in diluted Sulphuric Acid, and on evaporating
this solution an orange-red coloration is first produced, changing to a bluish-
violet and finally to a reddish-violet. Concentrated Sulphuric Acid dissolves
Narcotine with the production of a greenish-yellow colour, and on changing
to a yellow and finally to a reddish-yellow colour. On the addition of a trace of
Nitric Acid to its solution in concentrated Sulphuric Acid a beautiful red coloration
is produced, it is precipitated by the usual alkaloidal reagents, *e.g.*, Mercuric
Potassium Iodide (Mayer's) Solution, Iodo-Potassium Iodide (Wagner's) Solution,

PICRIC ACID, etc When heated with Nitric Acid it is oxidised with the formation of Cotarnine

Narcotine may be distinguished from the majority of other alkaloids by shaking with Acetic Acid Solution (2 p c) and filtering, the filtrate, when evaporated to dryness, should leave no weighable residue. It may be distinguished from Morphine by shaking with Sodium Hydroxide Solution (5 p c), the filtrate should yield no crystalline precipitate in 24 hours when treated with an excess of Ammonium Chloride Solution. It should leave no weighable residue when ignited with free access of air.

COTARNINÆ HYDROCHLORIDUM Cotarnine Hydrochloride, Stypticin, $C_{17}H_{15}NO_3 \cdot HCl$, eq 271.57. A pale yellow, crystalline powder, soluble in Water and in Alcohol. Cotarnine is produced by the oxidation of Narcotine, usually by means of Nitric Acid. Cotarnine Hydrochloride is the Hydrochloride of this oxidation product.

Dose — $\frac{1}{2}$ to $\frac{1}{2}$ grain = 0.021 to 0.032 gramme, given in capsule, or by hypodermic injection.

It may also be prescribed as a 1 in 10 Tincture, made with Tincture of Cinnamon. Dose, 10 drops in Water 4 times a day. — *B M J F* '01, ii 68.

Valuable in menorrhagia. Contra indicated in threatened abortion. — *P J* '95, ii 471, *B M J* '96, ii 17, *B M J E* '96, i 7, '98, i 71, 10.

In uterine hæmorrhage and in hæmorrhage during pregnancy, menopausal bleedings and post puerperal hæmorrhage. Failed almost entirely in all the cases of chronic metritis and endometritis. — *P J* Nov 41, *B M J F* '99, ii 86.

Principally useful in cases in which there is an unhealthy condition of uterine mucous membrane, but of little value in cases in which fibroid, cancer or other new growths are present, of very little value in connection with pregnancy, $2\frac{1}{2}$ grains is the minimum dose, repeated 3 or 4 times in the 24 hours, and continued over long periods of time. — *F T* '07, 15. One of the most valuable uterine hemostatics and sedatives. — *F T* '07, 90.

Tablets, each containing 0.05 gramme = $\frac{1}{2}$ grain, are made.

Tests — Cotarnine Hydrochloride dissolves readily in Water, forming a yellowish solution, and should be neutral towards Litmus. A 6 p c aqueous solution of the salt, when treated with Iodo Potassium Iodide Solution, yields a brownish precipitate, which re-crystallises from Alcohol and melts at about $142^{\circ}C$ ($287.6^{\circ}F$). If 3 drops of Sodium Hydroxide Solution (15 p c) be added to a solution of 0.1 of a gramme of the Hydrochloride in 3 c c of Water, the addition of each drop produces a milky turbidity which again disappears on shaking, the free base crystallising from the clear solution. The precipitate should be white, and the supernatant liquid clear and of a pale yellow colour. The crystalline base, when separated, should possess a m p of about $180^{\circ}C$ ($366^{\circ}F$), but the m p is stated to depend largely upon the rapidity with which it is heated, should the above supernatant liquid be turbid or strongly coloured, it indicates the presence of foreign impurities. The salt should leave no weighable residue when ignited with free access of air.

COTARNINÆ PHTHALAS (Styptol) — A micro crystalline powder, soluble in Water. Useful in arresting uterine hæmorrhage. May be given in doses of $\frac{1}{4}$ grain = 0.05 gramme, in powder or cachet. — *B M J E* '03, ii 96.

Three grains dissolved in 35 minims of Water can be used subcutaneously to rapidly arrest hæmorrhages. — *B M J* '05, i 311. Stated to possess the advantage over Stypticin of being less irritating. Employed as a 2 to 5 p c solution in superficial inflammations. For extensive eczema a 1 to 2 p c solution was found best, while a 5 p c solution was used for small furuncles. Internally it may be given in the form of powder in doses of 0.1 gramme ($\frac{1}{2}$ grain) or in the form of a tablet 0.05 gramme ($\frac{1}{4}$ grain) 4 to 6 a day on an average. — *M P* '05, ii 299.

Tests — When tested with solutions of Sodium Hydroxide Solution (15 p c) as described above under Cotarnine Hydrochloride it should yield a base possessing the m p of Cotarnine. It should leave no weighable residue when ignited with free access of air.

PAPAVÉRINA Papaverine $C_{15}H_{17}NO_4$, eq 336.66. — White, crystalline

needles, or colourless, trimetric prisms Insoluble in Water, sparingly soluble in Alcohol and Ether Strongly narcotic

Dose— $\frac{1}{2}$ to $\frac{1}{3}$ grain = 0.0054 to 0.0216 gramme

Tests—Papaverine melts at about 147°C (296.6°F) It yields when warmed with Sulphuric Acid a bluish-violet coloration, in Nitric Acid it dissolves with a dark red colour When treated with Chlorine Water it dissolves with a production of a greenish coloration, which on the addition of Ammonia Water changes after some time to a blackish-brown coloration It should yield no weighable residue when ignited with free access of air

Not Official

OREXIN.

CEDRARINE PHENYL-DIHYDRO-QUINAZOLINE

$\text{C}_{14}\text{H}_{12}\text{N}_2$, eq 206.62

A whitish amorphous powder, having a pungent taste, and having an irritating effect on the nostrils, inducing violent sneezing Insoluble in Water

Dose—1 to 5 grains = 0.06 to 0.32 gramme

OREXIN HYDROCHLORIDE ($\text{C}_{14}\text{H}_{12}\text{N}_2\text{HCl} \cdot 2\text{H}_2\text{O}$, eq 278.57)—In white needles, or as a white powder, soluble in Water and in Alcohol (90 p.c.), insoluble in Ether. It is said to possess a stimulating effect on the appetite, and to be found useful in the treatment of dyspepsia This salt is now entirely superseded by the Tannate

Dose.—2 to 8 grains = 0.13 to 0.52 gramme

OREXIN TANNATE—A pale yellow, amorphous, odourless and tasteless powder, insoluble in Water, 1 in 50 of Alcohol (90 p.c.) Introduced as a gastric tonic Useful in the anorexia of phthisis It has been recommended as a prophylactic against sea-sickness, and also to control the obstinate vomiting following Chloroform narcosis It is contra-indicated in hyperacidity of the stomach—*L* '00, 1, 1020, *B M J E* '02, 11, 96, *M A* '00, 493, *P J* '03, 1, 162

Dose—5 to 10 grains = 0.32 to 0.65 gramme, in a cachet, 1 or 2 hours before a meal It should not be prescribed with solutions containing Iron salts

Foreign Pharmacopœias—Official in Jap

Tests—Orexin Tannate when heated with Zinc dust or powdered Zinc it evolves a strong odour resembling Iso-nitile, and on treating this mixture with very dilute Hydrochloric Acid the filtrate yields a blue coloration on the addition of Chlorinated Lime Solution It should leave no weighable residue when ignited with free access of air

Not Official

OVI ALBUMEN

The liquid white of the Egg, *Gallus Bankiva* var *domesticus*, Temm, was official in the *B F* '85, and now appears in the Appendix of the *B P* '98 It is a glairy, viscid colourless, or pale yellowish liquid It may be obtained in the solid state by cautious evaporation at a temperature below 50°C (122°F) It is employed as an antidote in poisoning by Copper, Mercury, or Silver salts, and for certain purposes of clarification

Commercial dried Albumen is in thin, transparent flakes, which should be free from unpleasant taste or odour of putrefaction

It is coagulated by heat, and is then rendered white, opaque and insoluble, in this condition it is used as a test for Pepsin

By the action of the gastric juice or of Pepsin in weak Hydrochloric Acid solution, or by Hyposulphite, Albumen is first converted into acid Albumen or Syntonin, and finally into Peptone

A solution of Albumen is used as a means of proving the absence of Metaphosphoric Acid from Acetum Phosphoricum Concentratum.

Foreign Pharmacopœias—The dried white is official in Den, Dutch, Ger, Ital, Swed, the liquid white in the Fr, Mex and Port

EIGONS—Alpha and Beta Eigons are stated to be stable combinations of Iodine with Albumen and Peptone respectively, and the corresponding Brom-Eigons are similar preparations containing Bromine

α EIGON is a light, yellowish grey powder, possessing a faint odour Insoluble in Water, soluble in solution of Sodium Hydroxide forming Sodium α Eigon It contains about 20 p.c Iodine Employed internally and used as a dusting powder Introduced as a substitute for Iodoform and the Iodides, also used in veterinary practice —*L J '01*, 1 702

Dose—5 to 10 grains = 0 32 to 0 65 grammae

β EIGON—A light, yellowish brown powder, possessing a faint Peptone odour Soluble in Water, on which account, and as it is stated to be readily assimilable, it has been given in derangements of the stomach

The Bromine compounds have been employed as sedatives in doses of 10 to 15 grains = 0 65 to 1 gramme, 3 or 4 times daily —*B M J E '02*, 1 47

IODALBACID—A yellowish, tasteless, odourless powder, soluble in Water It is a combination of Iodine and Albumen, and is stated to be useful as a substitute for the alkaline Iodides

Dose—15 to 30 grains = 1 to 2 grammes,

OVI VITELLUS—The yolk of the Egg of *Gallus Banhna* var *domesticus* was official in *B P '85*, and now appears in the Appendix of the *L P '98* It is officially used in the preparation of Mixture Spiritus Vin Gallici, and is unofficially employed as an emulsifying agent

GLYCERITUM VITELLI—Fresh yolk of Egg 9 Glycerin 11, rub the yolk of Egg in a mortar with the Glycerin gradually added, and mix thoroughly —*U S P 1890*

LECITHIN (Choline Distearyl glycerophosphate) —A translucent, yellow or yellowish white, hygroscopic, waxy solid, which should be completely soluble in Chloroform It is a phosphorised organic constituent contained in considerable proportion in the yolk of Egg, from which it is chiefly prepared It has been employed in neurasthenia, in brain and nervous diseases, and in tuberculosis It may be injected hypodermically in doses of $\frac{1}{2}$ to 2 grains = 0 05 to 0 13 gramme, dissolved in sterilised oil —*L '02*, 1 392, 676, 687, 1119, *C D '01*, 11 725, '02, 11 155

Dose—1 to 5 grains given in the form of pill, granules, or as a confection

Lecitogen—A combination of Lecithin and Cocoa It occurs as a powder pleasant to the taste, is taken in doses of 3 or 4 teaspoonfuls daily in Milk or Water, and is useful in secondary anæmias —*B M J E '05*, 11 100

Not Official OXYGEN

A colourless, odourless, tasteless gas, it has been condensed to a liquid at a very low temperature and under great pressure, but as supplied for medical purposes it is in the form of compressed gas It may be prepared in small quantities by heating Potassium Chlorate mixed with half its weight of pure dry black Manganese Oxide, and subsequent purification of the gas, but on the commercial scale it is generally prepared from pure dry air by absorption with caustic Baryta

When employed medicinally, it is generally inhaled from bags connected with cast-iron cylinders, containing 10, 20 and 40 cubic feet of compressed gas, furnished with gun-metal taps

Ital requires that it shall be free from Carbonic Acid gas, from Chlorine compounds, and from Ozone

Medicinal Properties.—Useful in pneumonia, bronchitis, bronchial catarrh, asthma. It has also been employed in poisoning by coal-gas, and Carbon Monoxide. In the form of Hydrogen Peroxide it has been used in Cyanide poisoning.

A case of acute double pneumonia successfully treated with Oxygen —*L. 'J1*, ii 840

Case of fetid bronchorrhoea treated by inhalation for several hours daily —*B M J '02*, i 509

Treatment at the C T of 88 cases of various skin diseases, including also 9 cases of, these 50 were discharged cured, and 18 were greatly relieved, in all the cases of consumption the disease was arrested —*L. '03*, ii 274

Tried with success (*L. '05*, ii 686) in epileptic fits

In pneumonia is seldom required, and its value is stated (*B M J '05*, i 812, *L. '07* i 809) to be much over-estimated and disappointing

Its administration is stated (*B M J E '05*, ii 48) to have given favourable results in cases of chlorosis, more particularly those manifesting severe gastric disturbance and intolerant of Iron

Foreign Pharmacopœias—Official in Fr (*Oxygène*), Ital (*Ossigeno*); Mex and Span (*Oxygeno*)

OZONE—Is an allotropic modification of Oxygen, produced by passing a silent discharge of electricity through Oxygen Gas. This gas possesses a peculiar odour, somewhat suggestive of dilute Chlorine. It is a powerful oxidising agent. When present in the air in large quantities it frequently produces irritation of the mucous membrane

SODIUM PEROXIDE—A white amorphous powder, which dissolves in Water with a hissing noise, with evolution of heat and formation of Hydrogen Peroxide. It is a powerful oxidising agent.

Under the names of Biogen and Hopogan, Manganese Peroxide and Magnesium Peroxide have been prepared and introduced into commerce, they evolve Oxygen on contact with a dilute Acid

BENZOYL PEROXIDE—Well-formed white prisms, m p 108.5° C (218.3° F). Insoluble in Water, soluble in Oil to the extent of 2 to 3 p c. Prepared by the action of Sodium Peroxide on Benzoyl Chloride. It is a powerful disinfectant (*P J '05*, ii 330), useful in the treatment of burns, wounds and many skin diseases. It may be prescribed in oily solution or as the following ointment: Benzoyl Peroxide, 1, Vaseline, 5, Lanolin, 5

OXYMEL. *See* MEL.

OXYMEL SCILLÆ. *See* SCILLA.

Not Official.

PANCREAS ENZYMES.

Pancreatic Juice, the fluid secreted by the fresh and healthy pancreas of the pig, *Sus scrofa*, or of the ox, *Bos taurus*, is known to possess four distinct properties: (a) the conversion of proteids, (b) the conversion of Starch and Glycogen, (c) the emulsification of fats, and (d) the curdling of Milk. Each of these properties is attributable to a peculiar ferment or enzyme, which as originally present in the pancreas is in an insoluble and inactive condition, known as a Zymogen, the ultimate solution depending upon the conversion of this insoluble and inactive Zymogen into a soluble and active enzyme by the aid

probably of the digestive action of the intestinal ferment *Enterokina* e. The conversion may also be brought about by the action of diluted acids. The enzymes act only in neutral or alkaline solutions. Their action is suspended in feeble acid solutions, and when digested at 40° C (104° F) for an hour in a solution of Pepsin of the normal acidity of the stomach (equal to 0.2 p c Hydrochloric Acid), or when digested with gastric juice, they are destroyed. They are also destroyed in solution by heating to 75° C (167° F). The juice is precipitated by mineral acids, metallic salts, and by Tannic Acid. When treated with an excess of Alcohol (90 p c) or stronger, it is precipitated. It undergoes putrefactive change with great rapidity. The process of isolating the ferments in an active condition from the fresh juice of the pancreas gland, is extremely complicated and involves much time and careful work. The products usually obtainable commercially are of a widely divergent character, and, considered from the point of view of their proteolytic and amylolytic activity, some are quite inert.

An even greater divergence exists in solutions of the enzymes prepared for hypodermic use, than in the commercial dried ferments, some solutions indicating the devotion of but little attention in their preparation to the purposes which they are intended to serve.

Official Preparation.

LIQUOR PANCREATIS PANCREATIC SOLUTION

A liquid prepared from 5 oz. of the fresh fat-free pancreas of the pig, from which the external membrane has been removed, and finely divided by triturating with washed sand or powdered pumice-stone. The mixture is digested in a closed vessel in 20 fl oz. of Alcohol (20 p c) for a week and filtered.

Tests — A measured quantity of 2 c c of the solution is mixed with 0.2 of a gramme of Sodium Bicarbonate and 20 c c of Water, and the mixture added to 80 c c of fresh cow's Milk, previously brought to a temperature of 45° C (113° F). The mixture is kept at this temperature for 1 hour, at the end of this time the milk should be so completely peptonised that a portion removed and added to some Nitric Acid in a test-tube should no longer produce a coagulation. In carrying out the test it is preferable to add a little Ether to dissolve the fat, which may otherwise be mistaken for a coagulation.

The above resembles *Liquor Pancreaticus* (Benger), which was introduced in 1879.

Not Official

The ferments which have been prepared from the juice, and regarding which more or less definite knowledge exists at the present time, are **Trypsin**, **Amylopsin**, **Lipase** (Steapsin) and **Rennin** (Chymosin).

PANCREATINUM (Pancreatin) — Commercial Pancreatin is a mixture of the enzymes existing in the pancreas of the hog. It is not official in either the *BP* or the *PG*. The *USP* defines Pancreatin as a mixture of the enzymes naturally existing in the pancreas of warm blooded animals, usually obtained from the fresh pancreas of the hog (*Sus scrofa* var. *domesticus*, Gray), or the ox (*Bos taurus*, Linné), and consisting principally of Amylopsin, Myopsin, Trypsin and Steapsin, and proved to be capable, when assayed by the under-mentioned method, of converting not less than 25 times its own weight of Starch into substances soluble in Water. A method is also given of ascertaining its power of digesting soluble proteids.

It is a yellowish, cream coloured, or greyish amorphous powder, possessing a faint characteristic though not unpleasant odour, and a taste somewhat resembling meat. It dissolves slowly and almost completely in Water, but is only partially soluble in Alcohol (90 p c). It digests both soluble and insoluble Albumens, and when brought into contact with amylaceous material rapidly induces hydrolysis,

converting it into soluble products, *e.g.*, Sugar, Dextrose, or Maltose. It exhibits these powers to the greatest advantage in alkaline or in neutral solutions. The presence of mineral acids or an excess of alkali has a retarding influence upon its digestive activity, whilst, similarly to the juice, its activity is altogether destroyed by Pepsin in acid solution.

Foreign Pharmacopœias—It is official in Fr and Mex (Pancreatin), Ital (Pancreatina Medicinale), Jap (Pankreatinum), Span (Pancreatina), and US (Pancreatinum).

Tests—Pancreatin may be assayed for its proteolytic and amylolytic activity by its action upon the soluble Albumens of Milk and upon Starch. The *U.S.P.* employs a process of which the following embraces the essential features—A weighed quantity of 0.28 gramme of Pancreatin is mixed with 1.5 grammes of Sodium Bicarbonate and 100 c.c. of tepid Water. A measured quantity of 400 c.c. of fresh cow's Milk is raised to a temperature of 38° C (100.4° F), and the Pancreatin mixture is added, the whole being maintained at the above temperature for 30 minutes. At the end of this time the Milk should be so completely peptonised that a small portion when diluted with three times its volume of water should produce no coagulation when mixed with some Nitric Acid.

The *U.S.P.* method of determining the amylolytic activity is as follows—A weighed quantity of 7.5 grammes of Starch is mixed with 120 c.c. of Water and boiled until a translucent mixture results, which is then cooled to 40.5° C (105° F). A weighed quantity of 0.3 of a gramme of Pancreatin, dissolved in about 10 c.c. of Distilled Water at 40.5° C (105° F), is then added, the flask well shaken, and the temperature of the mixture maintained at 40.5° C (105° F) during 5 minutes, at the end of which time the Starch should have been converted into substances soluble in Water. 2 drops of Tenth-normal Volumetric Iodine Solution are mixed with 60 c.c. of Water, and 4 drops of the warmed converted Starch Solution are added to the mixture, either no coloration or at most a wine red colour should result, showing the presence of Dextrin and Maltose. The appearance of a blue colour will indicate the presence of unconverted Starch, and the Pancreatin is below the standard, namely, that of converting not less than 25 times its own weight of Starch into substances soluble in Water.

TRYPSIN—Trypsin is not official in the *B.P.*, *U.S.P.*, or *P.G.* It is given as a synonym for Pancreatin in the Spanish Pharmacopœia. It acts slowly on solid albuminoid masses, *e.g.*, boiled Egg Albumen, but with great rapidity on soluble Albumen, such as the Casein of Milk. It converts Albumens into Peptones and subsequently into bodies which are not proteids, Leucin, Tyrosin, etc. The activity gradually increases with the temperature up to 50° C (122° F), and rapidly diminishes up to 75° C (167° F) when the ferment is destroyed. Although the activity of the ferment is thus manifest with increasing temperature, solutions of the enzymes undergo rapid deterioration when subjected to prolonged warmth and light. Solutions, which have been subjected to temperatures such as 75° C (167° F) in a voyage through the tropics, have lost as much as 75 p.c. of their activity during their passage out and home.

Trypsin forms a yellowish or yellowish-brown powder, possessing a meaty odour. It occurs together with Amylopsin, Lipase (Steapsin), and Myopsin in the fresh juice of the pancreas. It is partially soluble in Water, insoluble in Alcohol (90 p.c.). The commercial dried product varies enormously, some specimens being exceedingly active proteolytically as well as amylolytically, others only relatively proteolytically active, whilst again other preparations possess neither a proteolytic nor an amylolytic activity.

Although Trypsin is not official in the *B.P.*, *U.S.P.*, or *P.G.*, a solution, *e.g.*, Sodium Bicarbonate or Sodium Carbonate, when in solution with the latter it rapidly deteriorates at a temperature of (or above) 38° C (100.4° F).

It is regarded (*J.C.S. Abs.* '05, 1147) as possible that the pancreatic enzyme Trypsin really consists of a number of specific ferments, each acting on different proteids. No free Trypsin is present in the secretion of the pancreas (*J.C.S. Abs.* '03, 11559), the liberation of that enzyme is the work of the testinal juice, gastric juice not being able to affect it.

Tests—Trypsin may be assayed for its proteolytic and amylolytic activity by the methods described under *Pancreatinum*, using either a proportionately smaller quantity of the enzyme or correspondingly reducing the time allowed for the reaction. The comparative strengths of the various Trypsin preparations are referred to under the heading of *Injectio Trypsini Co* (*Squac*).

AMYLOPSIN (Pancreatic Diastase)—It occurs together with Trypsin, Lipase (Steapsin), and Myopsin in the fresh juice of the pancreas. The dried ferment is not found in an active condition as a commercial article. Solutions of the ferment are extremely difficult to prepare, and even the best known products, although claiming to be free from the proteolytic ferment (Trypsin), can only be considered relatively so, as will be seen from the results published by Dr P. Tetenus Hald in his paper upon the comparative strengths of some commercial Trypsin preparations, in the *Lancet* '07, ii 1371. It possesses great activity in digesting amylaceous material, and when carefully prepared possesses but relatively little action on soluble or insoluble proteins.

Pancreatic diastase converts Starch into Dextrin and Maltose. Its action on Starch foods is very similar to or identical with that of Ptyalin, the salivary ferment. It is usually stated to be identical with the Diastase of Malt, but it is doubtful if it is so, as it is found to be affected quite differently to the latter by acid or alkali. Diastase from either source acts most rapidly in solutions which are practically neutral. The Malt ferment is retarded by acid, but almost stopped by a very small quantity (about 0.1 p.c.) of alkali. The pancreatic ferment, on the contrary, is retarded by alkali and almost stopped by a minute quantity of acid; moreover, the activity of Malt Diastase towards Starch Solution is inversely proportional to the quantity of the ferment present, whilst the activity of the Amylopsin (Pancreatic diastase) is inversely proportional to the square root of the quantity of ferment present.

Tests—The amylolytic activity of Amylopsin or solutions of Amylopsin may be determined by the test with Starch Solution described under *Pancreatinum*.

Medicinal Properties—The various pancreatic solutions, powders and tablets are used to peptonise foods previous to administration, but they are also given with food at the beginning of a meal. *Pancreatin* in pills (keratin coated) has been given in certain cases of diabetes. The enzymes of the pancreas gland have been employed with a certain measure of success in the treatment of malignant growths. A sufficient length of time has not elapsed, nor has a sufficient amount of evidence yet been accumulated, to enable a positive opinion to be expressed regarding their exact value in the treatment, but there is little doubt that they are now being given an extensive trial. There can also be little doubt that failure has in numerous instances been due to the use of almost inert preparations of the enzymes. That there is considerable variation in strength even in the best and most trustworthy preparations is evidenced by the comparative researches on the tryptic strength of different Trypsin preparations recorded (*L* '07, ii 1371) in the paper by P. Tetenus Hald.

The 'Problems of Cancer' formed the subject of a lecture at the Edinburgh Pathological Club, and the text appears in the *L* '04, ii 1200. A further communication was made in a lecture at the University of Liverpool on January 20th, 1905, and afterwards reported in the *J* '05, ii 283.

A record of the action on Jensen's mouse tumour is given in the *B M J* '06, i 140.

We (Editorial) certainly think that he (Morton) has made out a case for a trial on a larger scale—*B M J* '07, i 159. In the pathological reports and results of the microscopical examination of sections taken from the tumour mass of one of Morton's cases there were evidences of degenerative changes indicative of some destructive process, but whether they were due to X rays or to Trypsin it was impossible to say. No new nodules were observed from the time the patient received these large doses of Trypsin to the time of his second operation, nor did those already existing increase materially in size. Microscopic examination of the nodules showed the cancer was still active. There was no evidence that the Trypsin treatment had exerted any definite influence on the tumour cells—*B M J* '07, i 488, 520.

A paper on the Trypsin treatment for cancer appears in the archives of the

Middlesex Hospital (6th Cancer Report), from the observations the authors conclude that the course of cancer, both as a disease and as a morbid process, is unaltered by the administration of Trypsin and Amylopsin — *B M J* '07, 1 1447

The cell solution produced by Pancreatin is mostly only a circumscribed process, and when sufficiently large quantities of the ferment are injected there is no selection between the carcinomatous and healthy tissue. The pancreatic ferment possesses a theoretical interest only (Leyden and Bergell) — *Deut med Wochenschrift*, 1907, xxii 913, *B M J* '07, 1 161

Amongst a number of favourable conclusions to a report of experiments instituted by W J Morton to test Beard's statements, are the following — In all cases signs of amelioration in the progress of the disease have been observed, enlarged glands associated with cancer have rapidly diminished in size under the influence of Trypsin, Trypsin has a decided effect in reducing cancer cachexia and in improving the general health, patients frequently refer their greatest feeling of improvement to the period of time when they are taking Amylopsin followed by Trypsin, the pure diastase (Injectio Amylopsini) had much to do with the favourable results, Trypsin should be used in larger doses feeling one's way, for instance, from 20 to 30 minims daily for from 4 to 6 weeks and then resorting to Amylopsin, Trypsin deserves further trial — *B M J E* '07, 1 11

The direct action of Trypsin on growing cancer cells, as shown clinically and microscopically, is sufficient variant to continue the treatment in inoperable cases, especially in view of the fact that there are apparently no serious results that can occur from its use — *B M J E* '07, 1 27

Case of sarcoma of the testicle successfully treated with Trypsin in solution. At the end of the treatment the wound remained closed and the patient as far as one could feel, was completely absorbed, the lumbar pain had disappeared, and the man was able to follow his employment without fatigue — *B M J* '07, 1 79

A case of growing abdominal tumour treated by hypodermic injections of Trypsin and Amylopsin. The vomiting, nausea, and flatulence disappeared and the appetite improved, then gradually the pain lessened and the swelling also steadily diminished, while the weight regularly increased — *B M J* '07, 1 525

We (Editorial) have no wish to condemn the — we have heard of cases where improvement appeared to have — Trypsin injections. We have also heard of cases where no benefit occurred. When the treatment has been tested systematically both on mice and men it seems to have failed completely, and it is only by a thorough trial that we can come to a correct conclusion as to its value — *L* '07, 1 240

A record of six cases successfully treated by hypodermic injections of Trypsin, including a case twice operated upon at the Middlesex Hospital and finally discharged as hopeless — *G P*, 1907, 548

A case described as malignant disease of the cæcum treated by hypodermic injections of Trypsin. Whilst not wishing to minimise the beneficial effects which followed the injections and which began to be experienced almost immediately, it is not desired that the record of the case should raise false hopes — *G P* '07, 804

A case of extreme malignancy treated by hypodermic injection of Trypsin and local application of a pancreatic lotion. Decrease in size, disappearance of faecal, and discharge scarcely more than that from a simple healing ulcer — *G P* '07, 807

The technique of the Trypsin-Amylopsin injections — *G P* '07, 810, 818

A case of carcinoma of the liver treated by injections. Growth materially reduced in size, and nausea disappeared — *G P* '08, 178

Trypsin is of value in cancer. Carcinoma of the cervix uteri, rodent ulcer, and epithelioma are eminently suitable for the treatment. Gout, rheumatoid arthritis and chronic rheumatism ought to be treated by Trypsin injections when other better known methods have failed. Regarding Amylopsin when injected separately, in Trypsin, it has been found to be of no value in the treatment of cancer — *G P* '08, 1 80

INJECTIO TRYPSINI COMP (Squire) Squire's Compound Injection of Trypsin — A standardised sterilised liquid, prepared direct from the fresh and healthy pancreas of the pig. It is of maximum potency, containing a definite number of units of the proteolytic enzyme (Trypsin), and of the amylolytic

enzyme (Amylopsin) It is made in three strengths, known as Standards I, II, and III, and is contained in hermetically sealed glass capsules of a dark amber tint holding 1 cc, sufficient for an average dose of the injection

Dose—The average dose varies from 17 to 34 minims = 1.0 to 2.0 cc, subcutaneously injected deeply, not into the growth itself, but into the healthy tissue in the immediate neighbourhood of the growth, or into the back or buttock the injections being made daily and the dose gradually increased Small doses of Trypsin are quite useless

Method of Hypodermic Use—When required for use the glass capsule should be rinsed first in 1 in 1000 Corrosive Sublimate solution, then in sterilised Distilled Water, broken at the file mark on the neck, and the contents drawn into the carefully sterilised all glass hypodermic syringe The injections should be carried out under the strictest aseptic precautions, and the employment of heat avoided at all stages of the process

Injectio Trypsini Comp should be used at the commencement of the treatment, starting with 1 cc of Standard II, and if no bad symptoms arise, continue these injections daily for at least 6 weeks If after a few injections the patient gets lethargic or depressed, with headache and palpitation, put him for a few days on Standard I then on alternate days on Standards I and II, and subsequently, with careful watching, get back to Standard II every day After 2 months' treatment the medical practitioner must be guided by each individual case as to how much Trypsin and Amylopsin to give The method for the next month or so is to give about 5 Trypsins to 1 Amylopsin with 1 day's rest to each week After a long course of Trypsin, or if any septic symptoms arise, Amylopsin must be pushed, and even then the method is to give a Trypsin in the morning and an Amylopsin in the evening

INJECTIO AMYLOPSINI (Squire) Squire's Injection of Amylopsin—A standardised sterilised liquid of maximum potency, prepared direct from the fresh and healthy pancreas of the pig It contains a high number of Amylopsin units, and is relatively free from the proteolytic enzyme (Trypsin)

It is contained in hermetically sealed glass capsules of a dark amber tint holding 1 cc, sufficient for an average dose of the injection

Dose—The average dose varies from 17 to 34 minims = 1.0 to 2.0 cc, subcutaneously injected deeply, not into the growth itself, but into the healthy tissue in the immediate neighbourhood of the growth, or into the back or buttock, the injections being made daily and the dose gradually increased

Method of Hypodermic Use—When required for use the glass capsule should be rinsed first in a 1 in 1000 Mercury Perchloride solution, then in sterilised Distilled Water, broken at the file mark on the neck, and the contents drawn into the carefully sterilised all glass hypodermic syringe The injections should be carried out under the strictest aseptic precautions, and the employment of heat avoided at all stages of the process

The Amylopsin injection is meant to replace the Trypsin injection in the later periods of the treatment, and to meet bad symptoms, such as nausea, vomiting, pain in the back, drowsiness, albuminuria, etc, which may arise

LIQUOR PANCREATICUS FORT (Squire) Squire's Strong Pancreatic Solution—A standardised solution of the enzymes prepared from the fresh and healthy pancreas of the pig, for internal administration

Dose—1 to 2 fl drim = 3.6 to 7.1 cc 3 times daily half an hour before food

LIQUOR TRYPSINI COMP (Squire) Squire's Compound Trypsin Solution—A solution prepared on somewhat similar lines to the above

Dose—1 to 2 fl drim = 3.6 to 7.1 cc 3 times daily half an hour before food

TABELLÆ PANCREATICÆ FORT (Squire) Squire's Strong Pancreatic Tablets.—Tablets for internal exhibition, weighing about 8 grains, containing the digestive enzymes of the fresh and healthy pancreas of the pig

Dose—1 or 2 tablets 3 times daily half an hour before food

LOTIO PANCREATICA FORT (*Squire*) Squire's Strong Pancreatic Lotion—A standardised limpid liquid, containing the digestive enzymes of the fresh and healthy pancreas of the pig. It exerts a powerful solvent action on animal proteids, and is of value as a surgical solvent. For use as a pigment it may be applied locally with a brush undiluted, or when diluted 1 to 5 to 1 to 10 as a rectal or vaginal injection.

A corresponding Lotio Typhsim Comp (*Squire*) is also made.

GYNECOL AND ENICOL PANCREATICUS FORT (*Squire*) Squire's Strong Pancreatic Pessary and Suppository—These preparations are made for continuous local solvent action in such cases (uterine and rectal) as admit of this form of treatment.

They are made in small and large sizes.

LIQUOR PANCREATICUS—Glycerin of Pancreatin, 16.50, Sodium Bicarbonate, 3.50, Glycerin, 5, Alcohol, 15, Distilled Water, *q s* to produce 100—*B P C*.

ELIXIR PANCREATIN—Pancreatin, 5, Sodium Bicarbonate, 3, Alcohol, 15, Distilled Water, 45, Aromatic Elixir, *q s* to produce 100—*B P C*.

GLYCERINUM PANCREATIN *Syn* Glycerol of Pancreatin—Pancreatin, 10, Glycerin, 50, Simple Elixir, 5, Distilled Water, *q s* to produce 100—*P J F* and *B P C*.

PULVIS PANCREATICUS COMPOSITUS *Syn* Peptonising Powder—Pancreatin (*U S P*), 20, Sodium Bicarbonate, 80. Mix them by trituration—*U S N F*.

Note—1.5 grammes of this powder are sufficient to peptonise 300 c.c. of fresh cow's Milk or 25 grains will peptonise 20 fl. oz. in the following manner—Add 1.5 gramme of the Compound Pancreatic Powder to 125 c.c. of tepid Water contained in a suitable flask, and afterwards add 500 c.c. of fresh cow's Milk previously heated to 38° C (100.4° F). Maintain the mixture at this temperature for 30 minutes, then transfer to a cold place. Milk thus prepared should not be used after it has been kept for 24 hours or when it has developed a bitter taste—*U S N F*.

This has been incorporated in the *B P C*.

PULVIS PRO LACTE HUMANISATO *Syn* Humanising Milk Powder—Compound Pancreatic Powder (*N F*), 35, Sugar of Milk, 965. It is used for preparing Milk as follows—Triturate, 6.5 grammes of the Milk Powder with 62 c.c. of Water, transfer to a clean bottle containing 62 c.c. of fresh cow's Milk, and 15 c.c. of Fresh Sweet Cream, and immerse the bottle in Water heated to 38° C (100° F) for 15 minutes. Then pour the mixture into a vessel and heat it quickly to boiling and immediately allow it to cool to the body temperature. Humanised Milk should be prepared immediately before use and the directions carefully followed—*U S N F*.

This has been incorporated in the *B P C*.

PEPTONISED MILK—A pint of Milk is diluted with 4 fl. oz. of Water and heated to 140° F (60° C)*. To this add 2 teaspoonfuls of Liquor Pancreatis and 20 grains of Sodium Bicarbonate. Place in a jug and cover with a 'cosey' to keep it warm. At the end of an hour, or rather more, boil the contents of the jug. The product can be used like ordinary Milk.

Peptonised Milk can also be prepared at about 60° to 65° F. Dilute a pint of Milk with half a pint of Lime Water, or with half a pint of Water containing 20 grains of Sodium Bicarbonate in solution, to this add 3 teaspoonfuls of Liquor Pancreatis. The mixture is set aside in a jug for 3 or 4 hours, by which time the Milk will have developed a slightly bitter taste and will be ready for use.

The bitter taste is well covered by Soda Water, or it may be warmed and sweetened for infants.

* If a thermometer is not at hand, the proper temperature may be obtained by boiling one-half of the mixture and adding it to the other half which is cold,

If it is used as soon as ready it need not be boiled, but if not it must be boiled to prevent the change proceeding far enough to render it unpalatable.

Peptonising Powders and Tablets are also used in place of the Liquor Pancreatis. The powders generally contain the Sodium Bicarbonate mixed with the Pancreatin, ready for use.

PEPTONISED GRUEL - Gruel from wheaten flour, oatmeal, arrowroot, sago, pearl barley, pea or lentil flour, should be very well boiled and made thick and strong. It is then poured into a covered jug and allowed to cool to a lukewarm temperature. Liquor Pancreatis is then added, 2 teaspoonfuls to a pint of gruel. At the end of 3 hours the product is boiled and strained. The starch of the meal is converted into sugar, and the albuminoid matters are peptonised.

PEPTONISED MILK-GRUEL - To a good thick Gruel, prepared from any of the above mentioned farinaceous articles, while still hot, add an equal quantity of cold Milk, the mixture will be about 125° F (52° C). To each pint of this mixture add 2 teaspoonfuls of Liquor Pancreatis and 20 grains of Sodium Bicarbonate. Set aside in a warm place for 2 or 3 hours until a perceptible bitterness is developed and not longer, then heat to the boiling point and strain.

PEPTONISED BEEF-TEA - Half a pound of finely minced lean beef is mixed with a pint of Water and 20 grains of Sodium Bicarbonate. This is simmered for 2 hours in a covered saucepan. The resulting beef tea is decanted off into a covered jug, the undissolved beef residue is then beaten up with a spoon into a pulp and added to the Beef tea. When it has cooled down to about 140° F (60° C) a tablespoonful of the Liquor Pancreatis is stirred in. The mixture is kept warm for 2 or 3 hours and occasionally stirred. At the end of this time the contents of the jug are boiled briskly for 2 or 3 minutes and finally strained. Beef tea prepared in this way is rich in peptone, and when seasoned with salt is scarcely distinguishable in taste from ordinary beef-tea.

A concentrated preparation is supplied as **Peptonised Beef Jelly**.

PEPTONISED NUTRITIVE ENEMATA - The enema may be prepared in the usual way with milk gruel and beef tea, and a dessert-spoonful of Liquor Pancreatis should be added to it just before administration.

In the warm temperature of the bowel the ferments find a favourable medium for their action on the nutritive materials with which they are mixed.

It must be borne in mind that peptonised foods are very liable to change on keeping, and that fresh quantities should be prepared every 12 hours or they must be re-boiled. — *Sir W Roberts, Lumleian Lectures, 1880*

PANCREATISED FAT or PANCREATIC EMULSION

Introduced in the treatment of consumption and other wasting diseases, by Dobell.

Dose - From 1 to 4 fl drms = 3.6 to 14.2 cc, mixed in Milk or Water, from 1 to 4 times in 24 hours.

Not Official

PAPAIN.

Syn — PAPAYOTIN

A white, or whitish, amorphous powder, soluble in Glycerin. It is a digestive ferment extracted from Papaw Juice (*Carica Papaya*, L.)

Papaw leaves contain an alkaloid Carpaïne, the Hydrochloride of which is readily soluble in Water, it has been used as a heart tonic and febrifuge.

Papain possesses a solvent action on animal proteins, and acts best in neutral or slightly alkaline solution.

Some commercial Papains possess such activity in acid solution that they have been suspected of being admixtures containing Pepsin.

Medicinal Properties - Its solution (5 p.c.) is stated to dissolve false membrane in diphtheria, and to be a good application to warty epitheliomatous

growths—*B M J* '85, ii 151, '88, i 1296, *M P* '94, i 633, *P* h 372, *B M J E* '93, ii 39 Internally in gastric ulcer—*L* '94, i 840, '95, i 333 In atonic dyspepsia—*L* '95, i 1050 In gastritis—*B M J E* '93, ii 36

The results of a research (*L* '05, i 589) show that Papain contains a fibrin-digesting but not peptolytic protease of the nature of Pepain, as well as a peptolytic but not fibrin-digesting protease of the nature of an erepsin

Some further importance has recently been attached to this substance by its use in the treatment of malignant growths, attention being drawn to it by the publication of several letters relating to the action of certain other ferments on inoperable cancers A mass of scirrhous was injected (*B M J* '06, i 1439) with 2 grains of this substance Since then, the same procedure was adopted with three tumours of similar nature These injections were followed by a burning feeling of short duration and then by an occasional gnawing sensation The tumours softened in a few days, burst, and gave out a copious discharge of thick grey fluid for about a week When the discharge ceased, the lumps were found to have disappeared or become much smaller One tumour had to be injected 3 times before it burst The temperature rose from 2 to 4 degrees, but became normal in 2 or 3 days

Injections of $\frac{1}{2}$ grain and upwards into malignant growths with good results—*B M J* '07, i 185

Dose—2 to 10 grains = 0.13 to 0.65 gramme

Prescribing Notes—*May be given in cachets, mixture, pills, or as a hypodermic injection A good pill may be made by using 'Dispensing Syrup' q s Given also in the forms of Elixir and Glycerole, in doses of 1 teaspoonful*

Foreign Pharmacopœias—Official in Mex

ELIXIR PAPAIN (*Squire*)—Glycerinum Papain (*Squire*), 8 fl oz, Carmine solution, 2 fl drim, Spiritus Nucis Juglandis, 2 fl drim, Elixir Aurant, sufficient to produce 16 fl oz

1 fl drim (8.6 cc) contains $2\frac{1}{2}$ grains (0.16 gramme) of the purified and dried juice

Dose—1 to 2 fl drim = 3.6 to 7.1 cc twice or thrice daily, half an hour before food

Papain, 11, Saccharin, 0.4, Glycerin, 60, Sherry, 150, Chloroform Water (1 in 200), 890—*Hager*

Papain, 5, Alcohol, 15, Distilled Water, 45, Aromatic Elixir, q s to produce 100—*B P C*

GLYCERINUM PAPAIN (*Squire*)—Papain purified and dried, 640 grains, Sodium Bicarbonate powder, 40 grains, Glycerin, 8 fl oz, Aqua Dest sufficient to produce 16 fl oz

Papain, 1 oz, Hydrochloric Acid, 40 minims, Purified Talc, 120 grains, Glycerin, 8 fl oz, Water, to 16 fl oz—*Pharm Form*

Papain 8 Diluted Hydrochloric Acid, 8, Simple Elixir, 5, Glycerin, q s to produce 100—*B P C*

INJECTIO PAPAIN FORT (*Squire*)—A sterilised limpid liquid possessing the full digestive powers of the purified and dried juice 1 cc (17 minims) contains 2 grains (0.13 gramme) of the purified and dried juice

Dose—1 to 2 cc = 17 to 34 minims, hypodermically injected deeply into the subcutaneous tissue

The above injection is prepared in the form of hermetically sealed glass capsules of a dark amber tint, each capsule containing sufficient for an average hypodermic dose

LOTIO PAPAIN (*Squire*)—A clear permanent solution of the enzymes in a Glycerin basis, prepared for use as a surgical solvent It may be applied locally undiluted, with a camel's-hair brush

PAPAPERIS CAPSULÆ.**POPPY CAPSULES**

FR, PAVOT, GER, UNREIFE MORHAKOPFF, ITAL PAPAVERO,
SPAN, ADORNIDIRA

The nearly ripe dried Fruits of the Opium Poppy, *Papaver somniferum*, L

Medicinal Properties Similar to Opium, but much weaker and of uncertain strength The decoction is used as a soothing anodyne fomentation

Not Official -Decoctum Papaveris, Sympus Papaveris and Extractum Papaveris Liquidum

Foreign Pharmacopœias Official in Austr, Belg, Din, Dutch, FF- (Pavot), Ger, Hung, Mex (Adornideras), Port (Dormideras), Russ, Span (Adornidera), and Swiss Not in the others

Descriptive Notes Poppy capsules are usually dried gradually on the plant, the stalk being bent downwards as soon as the poppy head has arrived at its full size, and the capsule is thus allowed to become hard on the plant The first capsule formed is usually the largest, the subsequent, smaller, capsules are sorted out, and sold separately, and the smallest are usually broken up and sold at a lower price, for making fomentations, etc There are two forms of the capsules, viz (1) nearly spherical, and depressed at base and apex, and (2) oblong oval The seeds are white when derived from the white flowered form, but are often greyish when derived from the red flowered variety of *Papaver somniferum* The white seeded form is the one official in the *BP* The capsule is described as being the nearly ripe dried fruit usually 2 to 3 in (5 to 7½ cm) in diameter, and suddenly contracted below into a neck, and covered above with stellately arranged stigmas, the pericarp being pale yellowish brown externally, and frequently marked with dark spots, and having a bitter taste but no odour The seeds, and the fixed Oil derived from them, are both official in the *PG*, in which the unripe capsules are directed to be cut in half longitudinally and freed from the seeds before use, without which they should weigh 3 to 4 grammes and show dried milky juice at the edges The seeds are described as reniform, 1 mm long, with a network of six sided meshes on the surface Neither the seeds nor the capsules nor the Oil are official in the *USP*

Tests. - Poppy capsules yield about 10 p.c. of ash

Not Official

DECOCTUM PAPAVERIS -Poppy Capsules, bruised, 2, Distilled Water, 80, boil 10 minutes in a covered vessel, and strain, then pour over the contents of the strainer as much Distilled Water as will make up the strained product to 20 (1 in 10)

An external soothing application, applied warm

Foreign Pharmacopœias. -Span, Infusion, 1 in 35

SYRUPUS PAPAVERIS (*BP* '85) -3G of Poppy Capsules is exhausted with boiling Water, and the liquid evaporated to 60, this is treated with 1G of

Alcohol (90 p c), and subsequently evaporated to 40, in which is dissolved 64 of Sugar (1 in nearly 2½)

Dose—1 fl drim = 3 6 c c

On the average, 60 minims will equal 8 minims of Tincture of Opium

This is made from the Liquid Extract (*given below*) by evaporating 80 of the to 40, and dissolving in it 64 of Sugar

Liquid Extract (*given below*), 40, Sugar, 70, dissolve and make up to 100—*B P C*

This is less than half the strength of the preparation given above, but the difference is probably unintentional

Foreign Pharmacopœias—Official in Austr (Syrupus Opiatus)—Extract Opium 1, Simple Syrup 999 Dutch, Ger and Russ, 1 in 10, Belg (Syrupus Opii dilutus), Syrup of Opium 1, Simple Syrup 4 Dan, about 1 in 12, Hung (Syr Diacodii) 1 in 27, Ital (Sciroppo di Opiio), Tinct of Opium, 1 in 1000, Mex (Jarabe diacodio), 1 of Ext Opium in 2000, Port (Xarope de Dormideiras), 1 in 13½, Span (Jarabe de Adormideiras), 1 extract in 100, Swiss, Extract of Opium 1, Water 4, Simple Syrup 999 Not in Fl, Ital, Jap, Norw, Swed or U S

EXTRACTUM PAPAVERIS LIQUIDUM—The liquid obtained by the process for making the Syrup (previous to adding the Alcohol and the Sugar), 8, Alcohol (90 p c), 1, mix

Dose—30 to 60 minims = 1 8 to 3 6 c c

This has been incorporated in the *B P C*

PARAFFINUM.

Petroleum Oil and Shale are mixtures of the hydrocarbons of the Paraffin series, some of which are official under the names Paraffinum Durum, Paraffinum Liquidum, Paraffinum Molle Hard Paraffin is obtained chiefly from Shale, the Liquid and Soft Paraffins from Petroleum

PARAFFINUM DURUM. HARD PARAFFIN

Fl, Paraffine, Ger, Festes Paraffin, Ital, Paraffina, Span, Paraffina

A colourless, crystalline, wax-like solid, which is a mixture of several of the harder members of the Paraffin series

Solubility—Insoluble in Water, soluble in Absolute Alcohol, 1 in 80 of Ether, sp gr 0.720, 1 in 40 of Ether, *B P*

In *B P* '85 it was stated to be 'freely soluble in Ether, which is altered in *B P* '98 to 'almost entirely soluble in Ether'

The solubility in Ether (sp gr 0.720) depends upon the m p of the Paraffin, a sample m p 120° F, dissolved 1 in 40

Official Preparation—Unguentum Paraffini Contained in Unguentum Crocosi and Unguentum Calaphiti

Not Official—Emulso Paraffini and Massa Paraffinum

Foreign Pharmacopœias—Official in Belg, Ger, Hung, Jap and Russ, all Paraffinum Solidum (m p 74° to 80° C), Dutch m p 56° to 60°, Fr, Paraffine (distils between 375° and 435°), m p not given, Span, m p 44° to 65°, Swiss, m p 65° to 80°, U S, Petrolatum Spissum (m p 45° to 48° C) Not in the others

Paraffin Injections—Very frequent references to this method of remedying deformities have been in recent years. An exhaustive paper (*B M J* '04, ii 1154) shows that the best form of syringe is an all-metal one with a screw piston. The

Paraffin is melted by placing the containing bottle up to it neck in very hot Water. Then it and the syringe are kept in Water about 2' above the m.p. of the Paraffin. The syringe is filled, the needle dipped for a few minutes in very hot Water to prevent the Paraffin from setting in the needle, the needle is inserted well under the skin and the injection made quickly.

Thrombosis is avoided by using screw syringe, so that Paraffin may be injected slowly and steadily.

Ethyl Chloride should not be used to cool the Paraffin, but only a stream of cold Water. The Paraffin used should have a m.p. about 10° F and the injection should be made about 120° F. About 1 cc may be enough in case of sunken nose. In prolapse of the rectum or the vagina 1 oz, or in bad cases 1½ oz, may be required (*B.M.F.* 04, ii 1155).

For remedying deformities of the nose an interesting method of procedure is given in the *L.M.F.* 04, ii 1239. The Paraffin used is a mixture of Hard and Soft Paraffin having m.p. of 106° F. From an experience of over one hundred cases it has been shown that the operation, if properly performed is practically devoid of danger.

For external injections a Paraffin liquefying between 45° and 50° C (113° and 131° F) should be used (*B.M.F.* 06, i 1108) while for the treatment of atrophic *osena* cold injection of Paraffin melting at 13° C (113° F) should be used.

Some cases of bone cavities treated by stopping with Paraffin method described, Paraffin of a m.p. of 120° F should be used, *L.* 08, i 155.

A new syringe, *L.* '07, i 1612.

Tests—Hard Paraffin has s.p.g. according to the *B.P.* from 0.820 to 0.940, the *U.S.P.* gives the s.p.g. at 0.890 to 0.905 at 25° C (77° F). It melts according to the *B.P.* at 54.4° to 57.2° C (130° to 135° F). The *U.S.P.* m.p. is 51.6° to 57.2° C (125° to 135° F). The *P.G.* m.p. is 74° to 80° C (165.2° to 176° F). No s.p.g. figure is included in the *P.G.* The *B.P.* states that it melts at the temperature given above, and burns with a bright flame. The *U.S.P.* more correctly states that when strongly heated it ignites and burns with a luminous flame. The *U.S.P.* states that if 0.5 of a gramme of Paraffin be heated in a dry test-tube with 0.5 of a gramme of Sulphur, Carbon is separated and the mixture becomes black, Hydrogen Sulphide gas being simultaneously evolved, no similar test is given in either the *B.P.* or the *P.G.*, all three Pharmacopœias state that in alcoholic Solution should not redden blue Litmus paper.

The more generally occurring impurities are free acid, Stearic Acid, and fixed Oils. The presence of free acid may be shown by the acid reaction of the alcoholic solution, fixed Oils or other organic impurities may be shown by the Sulphuric Acid test described below, and the presence of Stearic Acid by the Fuchsin test described below, and also by the acid nature of the solution in Alcohol (90 p.c. or 94.9 p.c.). It should leave no weighable residue when ignited with free access of air.

Sulphuric Acid—It should not be acted upon nor coloured by Sulphuric Acid, *U.S.P.* 3 grammes heated on a water bath with 3 cc. of Sulphuric Acid in a glass previously rinsed out with warm Sulphuric Acid and carefully agitated for 10 minutes should not be affected and the acid should not be coloured more than faintly brown, *P.G.*

Nitric Acid—It should not be acted upon nor coloured by Nitric Acid, *U.S.P.*

Fuchsin.—If 0.5 gramme of Paraffin and 0.1 gramme of powdered Fuchsin added to the fused mass, the latter should not assume a pink or red colour, *U.S.P.*

Water to produce 24 fl oz.—*B P C Formulary* 1901, now incorporated in the *B P C*, adding 1 p c by volume of Elixir Glusidi

OLEUM PRO NEBULA—Purified White Petroleum Oil, 1 fl oz, Balsam of Peru, 40 grains Digest in a bottle on a water-bath for 10 minutes, and filter when cold—*Bournemouth Formulary*

PARAFFINUM MOLLE.—SOFT PARAFFIN

A semi-solid translucent substance Either the white or the yellow variety may be used, according to circumstances

Vaseline, Adepsine, Salvo Petrolia, Chrisma, and Cosmoline are forms of Soft Paraffin

Solubility.—Insoluble in Water, slightly soluble in Absolute Alcohol, freely in Ether, Chloroform, Benzol, Oil of Turpentine, the fixed and volatile Oils

Description of a syringe suitable for making hypodermic injections—*L '03*, 11 611, *B M J '03*, 11 741

Official Preparations—Unguentum Paraffini The White is used in the preparation of *Unguentum Eucalypti* and *Unguentum Zinci Oleatis* The Yellow in *Unguentum Hydrargyri Nitratis Dilutum* and *Unguentum Hydrargyri Oxidi Flavi*

Not Official—Emulso Paraffini, Massa Paraffinum, Ceratum Paraffini, Linogenum Spissum, Parenols, Parogens, Vasenol, Vasogen, and Vasolmenta, Petroleum Spirit (Petroleum Ether)

Foreign Pharmacopœias—Official in Austr, Dan, Jap, Norw, Swed and Swiss (Vaselinum), Belg (Paraffina Mollis), Dutch (Vaselinum Album and V Flavum), Fr (Vaseline Officinale), Ger and Russ (Unguentum Paraffini), Hung, Ital and Span (Vaselina), Mex (Vaselina Solida), US (Petrolatum Molle)

Tests.—Soft Paraffin is officially stated to possess a sp gr of 0.840 to 0.870 at the temperature of its m p The *USP* states that it has a sp gr of from 0.820 to 0.850 at a temperature of 60° C (140° F) The m p is given in the *BP* as from 35.5° to 38.9° C (96° to 102° F) or even somewhat higher The *USP* gives the m p as between 45° and 48° C (113° and 118.4° F)

The more generally occurring impurities are free acid, readily carbonisable impurities, fixed Oils, fats, Rosin, and mineral matter Soft is digested with Alcohol (90 p c), and the insoluble oily matter separated, the alcoholic solution should be neutral in reaction towards Litmus paper When mixed with twice its volume of Sulphuric Acid and warmed in a water-bath for 15 minutes the acid should not be coloured more than a light brown, when boiled with Sodium Hydroxide Solution, and the aqueous alkaline liquid separated from the oily residue, should yield no precipitate or oily matter on acidifying with Sulphuric Acid The *USP* uses the same test as is described under Liquid Paraffin A portion of the sample when carefully ignited with free access of air should leave no weighable residue Paraffin Molle is not official in the *PG*.

Litmus—Water shaken with melted Petrolatum should not redden blue Litmus, *USP*

Sodium Hydroxide.—Digest 10 grammes of Soft Paraffin, 10 grammes of Sodium Hydroxide, and 50 c c of Water for half an hour on a water bath and

then separate the aqueous layer. No oily or solid substance should separate from this when it is supersaturated with Sulphuric Acid, *U S P*

Sulphuric Acid—When 2 volumes of conc. Sulphuric Acid and 1 volume of melted Petrolatum in a test tube be placed in hot Water for 15 minutes, with occasional agitation, the acid should not acquire a deeper tint than brown nor lose its transparency, *U S P* 1906, but deleted in list of Additions and Corrections 1907

Not Official

EMULSIO PARAFFINI *Syn* Aseptic Shaving Cream.—Hard Paraffin (m.p. 55° C), 22, Prepared Suet, 3, Soft Soap, 2, Tragacanth, in powder, 2, Glycerin, 2, Oil of Lavender, 1, Boiling Water, 68. Place the Hard Paraffin and Suet in a vessel surrounded by hot Water, add the Soap and boiling Water and vigorously beat the mixture until a smooth white emulsion is obtained. Remove the surrounding hot Water, and gradually add the Tragacanth, continuing the beating and stirring until the temperature has fallen below 50° C. When nearly cold, add the Glycerin and Oil of Lavender. This product should have the consistence of a soft paste, and is used to facilitate the shaving of skin areas so as to obviate the use of a shaving brush and soap.—*St. Thomas's*

This has been incorporated in the *B P C*

MASSA PARAFFINUM—Hard Paraffin (m.p. 120° F), 1, White Soft Paraffin 1½, melt together

A good mass for making Silver Nitrate and Potassium Permanganate into Pills

This has been incorporated in the *B P C*

Ceratum Paraffini—Beeswax, 6, Soft Paraffin, 94.—*B P C*

PARENOL—It has been shown by A. Kopp (*Apotheker Zeitung*, 19, 786) that Soft and Liquid Paraffin can be formed into stable emulsions with Water by the addition of a small quantity of Wool Fat, Beeswax, Spermaceti or other substances, consisting chiefly of the higher Alcohols or esters of those Alcohols. The resulting emulsions are absorbed readily through the skin, cause no irritation, and do not become rancid, while they serve a useful purpose as vehicles for the application of various medicaments.

The following formulæ are found to yield satisfactory products, the first being somewhat better than the second and third.

Wool Fat Parenol—Soft Paraffin, 65, Wool Fat, 15, Distilled Water, sufficient to produce 100. Warm the Water, and mix gradually with the melted Soft Paraffin and Wool Fat in a warm mortar.

This has been incorporated in the *B P C*

Beeswax Parenol—Soft Paraffin, 70, white Beeswax, 5, Distilled Water, sufficient to produce 100. Proceed as in the former case.

Spermaceti Parenol—Soft Paraffin, 70, Spermaceti, 5, Distilled Water, sufficient to produce 100. Proceed as in the first case.

These solid Parenols are of ointment-like consistence, can be made to take up more than their own weight of Water, mix with all fats, and can be used alone or in combination with other substances.

Liquid Parenol—Liquid Paraffin, 70, white Beeswax, 5, Distilled Water, sufficient to produce 100. Proceed as in the case of Wool Fat Parenol.

This has been incorporated in the *B P C*

The Liquid Parenol is a neutral liniment, possessing similar properties to the solid preparations, and can be used in the treatment of skin diseases, for lubricating catheters, or as a vehicle for injections.—*P J '06*, ii 623, *Y B P '07*, 278

VASOGEN (Oxygenated Vaseline Valsol)—A yellow, or dark brown, thick oily liquid which forms with Water a stable white emulsion. It has been introduced as a basis for various medicated preparations, *e g*, Creosote Vasogen (5 and 20 p.c.), Ichthyol-Vasogen (1 p.c.), Iodene Vasogen (6 and 10 p.c.), Iodoform Vasogen (15 p.c.), and Menthol Vasogen (2 p.c.)

A mixture of Vaseline Oil and Oleic Acid, when saturated with ammonia,

yields a similar preparation to Vasogen—*Proc Amer Pharm Assoc* xliii 632; *P J '02*, ii 259

A brown fatty solid is also known under the name of **Vasogenum Spissum**

VASENOL—A yellow vaseline preparation containing 25 p c Water, introduced as an ointment basis, and a liquid **vasenol** (white), which may be medicated as desired, has also been introduced where a creamy application is used—*B M J '04*, ii 1414

A combination of a Liquid Paraffin with a small quantity of the higher Alcohols obtained from Spermaceti, Wool Fat, etc. The product mixes readily with aqueous liquids producing neutral emulsions—*L '05*, i 1396

VASOLIMENTA—Under this name (*Pharm Cent*, xli 756) a combined soap hydrocarbon basis for medical injection has been introduced *Simple Vasoliment* or *Liquid Vasoliment*, is prepared by saponifying Oleic Acid, 50, with Alcoholic Ammonia, 25, the soap being heated with Liquid Paraffin, 100, until solution is effected. The weight is then made up to 175 with Alcohol. *Thick Vasoliment* is prepared in a similar manner, but the Alcohol is evaporated off. Medicated Vasoliments are prepared as solutions of the active ingredients in simple Vasoliment in the following percentage proportions respectively: *Salicylic Acid*, 2 p c, *Chloroform*, Camphor 30 p c and Chloroform 30 p c, *Iodine*, 6 p c, *Creolin*, 5 p c, *Menthol*, 2 p c, *Venice Turpentine*, 20 p c, *Iodoform*, 15 p c, *deodorised Iodoform*, Iodoform, 15 p c, *Eucalyptol*, 15 p c, *Eucalyptol*, 20 p c, *Naphthol*, 10 p c, *Guaiacol*, 20 p c, *Thiol*, 5 p c—*Y B P '01*, 212 and *Hager*

PAROGENUM *Syn* Liquid Parogen, Vasoliment, Oxygenated Paraffin—Liquid Paraffin, 40, Oleic Acid, 40, Ammoniated Alcohol (5 p c), 20—*B P C*

PAROGENUM SPISSUM *Syn* Thick Vasoliment—Hard Paraffin, 12, Liquid Paraffin, 48, Oleic Acid, 30, Ammoniated Alcohol (10 p c), 10, evaporate to 90 by weight—*B P C*

Vasoliments have been incorporated in the *B P C* under the title **Parogeni**

Mindes' Vasoliments—No 1 Liquid Paraffin, 35, White Olein, 35, Alcoholic Solution of Ammonia, 25, strong Alcohol, 5. No 2 Liquid Paraffin, 35, white Olein, 35, Alcoholic solution of Ammonia, 30. The No 2 formula is recommended for the preparation of a solution of Iodine, or of any other medication soluble in Ether

Linogens are obtained by substituting Linseed Oil for Liquid Paraffin in the two preceding formulæ

Linogenum Spissum—Linseed Ointment (Linseed Oil, 3, Paraffin, 2), 60, white Olein, 30, Alcoholic solution of Ammonia, 10. Mix intimately in a mortar. The product, of a bright, yellow colour, readily absorbs large quantities of water

Linogens of Iodine, Creolin, Creosotal, Creosote, Ichthyol, Resorcin and Veratrine are obtained by simple solution of the prescribed quantities of the active ingredient in liquid Linogen. Iodine Linogen containing 6 to 10 p c of Iodine, although quite bright when first made, becomes cloudy on keeping, especially if exposed to light—*P J '02*, ii 415

PETROLEUM SPIRIT *Syn* PETROLEUM ETHER—Now appears in the Appendix of the *B P '98*, and is there described as 'a colourless, very volatile and highly inflammable liquid. Sp gr 0.670 to 0.700, boiling point 120° to 140° F.' and is used as a solvent, the Petroleum Ether for use in conjunction with Methylated Ether (sp gr 0.717) for the production of local anaesthesia, has a much lower sp gr (0.640), and boils at a much lower temperature.

PARALDEHYDUM.

PARALDEHYDE

 $C_6H_{12}O_3$, eq 131 10FR, PARALDI HYDE, GER, PARALDI HYD, ITAL, PARALDI IDIF,
SPAN, PARALDI HIDO

A colourless, transparent mobile liquid having a peculiar characteristic, not unpleasant ethereal odour, and a pungent and subsequently a cooling taste. A polymer of Acetaldehyde.

It should be kept in amber-coloured stoppered bottles, and in a cool atmosphere.

Paraldehyde not answering the official requirements can generally be brought up to the standard by washing with Water containing an excess of Sodium Bicarbonate to remove acidity, and then dehydrating over dried Potassium Carbonate. If the m p be very low it should first be redistilled and the first tenth rejected.

Solubility—1 in $8\frac{1}{2}$ of Water at 60° F, the solution becoming very turbid on warming. It is miscible, in all proportions, with Alcohol (90 p c) and with Ether.

Medicinal Properties—Hypnotic. Produces quiet and refreshing sleep more speedily than Chloral, does not depress the heart's action. Has a marked action on the kidneys, increasing the flow of urine. It does not give rise to headache. Is a valuable remedy in the insomnia of cardiac disease, of mania, melancholia, and of other mental diseases.

Paraldehyde is given off by the lungs, and may be detected in the breath 12 or more hours after it has been taken.

30-minim doses every half or one hour in spasmodic asthma—*B M J* '93, 1 65, '96, 1 725, *L* '99, 1 756.

In 1 to $1\frac{1}{2}$ fl drms doses, one of the most potent remedies in spasmodic asthma—*Scot Med and Surg Jour* '99, 418.

One of the best and safest drugs for use as a narcotic in the treatment of mental diseases—*L* '02, 1 1539.

Cases of habit have been recorded. Over doses have occasionally produced epileptiform convulsions, but muscular relaxation and deep stupor are more frequent—*L* '03, 1 1023.

Sometimes useful in the sleeplessness of patients suffering from Bright's disease—*Pr* lxvii 658.

Case of poisoning by 1 oz of pure Paraldehyde given in mistake for a diluted preparation, recovery—*L* '02, 11 673.

Has been successfully employed in the insomnia of tricuspid incompetency—*M P* '04, 11 515.

A very good hypnotic—*B M J* '05, 11 1007. $1\frac{1}{2}$ drms may be given in 2 oz of Water, the disagreeable taste being covered by Tincture and Syrup of Orange. It is particularly valuable when insomnia is associated with delirium or any mental aberration.

Of special service as a hypnotic in chronic alcoholism—*B M J* '05, 11 250.

In severe bronchopneumonia and capillary bronchitis of infants, with Potassium Iodide and Liquorice—*B M J* '08, 1 258.

Dose.— $\frac{1}{2}$ to 2 fl drms = 1 8 to 7 1 c c

Ph Ger maximum single dose, 5 0 grammes, maximum daily dose, 10 0 grammes.

Prescribing Notes—May be taken dissolved in 1 to 2 fl oz of Water. A small dose repeated in an hour is more effective than a large dose. It is very pungent, when prescribed in mixture it should be diluted 1 to 16 of Water. The flavour is disagreeable and difficult to cover, Tincture of Orange and Cinnamon Water are the best for this purpose. When larger doses than will dissolve are required in mixtures, Compound Tragacanth Powder should be ordered to diffuse it. It is also prescribed in capsules.

It has been stated (*P J* '01, 1 559) that when Potassium Bromide and Paraldehyde are prescribed together in Water, that Potassium Bromate is formed, but we dissolved 3 grammes of Potassium Bromide and 4 grammes of Paraldehyde in 150 grammes of Distilled Water, and on keeping over a month the Bromide titrated its full strength.

Not Official—Metaldehyde, and Mistura Paraldehydi.

Foreign Pharmacopœias—Official in Dan, Hung, Ger, Ital, Jap, Mex (Paraldehyde), Norw, Russ, Span, Swiss and U S. Not in the others.

Tests—Paraldehyde has a sp gr of 0.998 to 0.999. The *BP* gravity is 0.998. A carefully fractionated sample of Paraldehyde may have a sp gr of 0.999. The *USP* sp gr is 0.990 at 25° C (77° F). The *PG* states 0.995 to 0.998. A carefully fractionated sample boils at 125° to 126° C (257° to 258° F). The *BP* boiling point is 124° C (255.2° F). The *USP* gives 121° to 125° C (249.8° to 257° F) and the *PG* gives 123° to 125° C (253.4° to 257° F) as the boiling point. It solidifies to a crystalline mass when cooled to a temperature of about 0° C (32° F) and melts again at 10° C (50° F). The *BP* gives the mp as 10° C (50° F), the *USP* as 10.5° C (51° F), and the *PG* as 10.5° C (51° F), but not under 10° C (50° F). The mp of a carefully fractionated sample is from 12.2° to 12.8° C (54° to 55° F).

It is neutral or but faintly acid in reaction towards Litmus paper. When warmed with Silver Ammonio-nitrate Solution the silver is reduced and on standing forms a metallic mirror on the sides of the tube.

The more generally occurring impurities are Aldehyde, Sulphates, Chlorides, free acid and impurities derived from Fusel oil. The *BP* has a test for the absence of Aldehyde, requiring that no coloration shall be yielded when the Paraldehyde is mixed with Potassium Hydroxide Solution and allowed to stand for 2 hours. The Aldehyde reaction with Potassium Hydroxide Solution is an exceedingly delicate one, almost too delicate, very few samples remaining quite uncoloured for 2 hours. No similar test is included in the *USP* or the *PG*. 1 c.c. of the specimen should form a clear solution with 10 times its volume of Water, which should be free from oily drops (absence of Amyl Alcohol), should yield no turbidity or precipitate with Barium Chloride Solution (absence of Sulphates), when acidified with Nitric Acid should yield no turbidity or precipitate with Silver Nitrate Solution (absence of Chlorides). The absence of any disagreeable odour when the Paraldehyde is carefully and completely evaporated indicates the absence of impurities derived from Fusel oil. The amount of free acid in the sample may be judged by the reaction towards Litmus. The *USP* and the *PG* both give a limit for free acid, the former requiring that a mixture of 8 c.c. of Paraldehyde and 8 c.c. of Alcohol (94.9 p.c.) should acquire a pink colour upon the

addition of 0.5 cc of Normal Volumetric Potassium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, the latter that a mixture of 1 cc each, of Paraldehyde and Alcohol (90 p c), shall not possess an acid reaction after the addition of 1 drop of Normal Volumetric Potassium Hydroxide Solution, but no indicator of neutrality is mentioned. Paraldehyde should be readily and completely volatilised at a temperature of a water-bath, and should leave no weighable residue.

Not Official

MISTURA PARALDEHYDI—Paraldehyde, 1 fl drim, Glycerin, 40 minims, Rectified Spirit, 2 fl drim, Cinnamon Water, to 1 fl oz.

Metalddehyde, which is also a polymer of Ethylic Aldehyde, occurs in colourless acicular crystals, and was at one time said to be a hypnotic in doses of 2 to 8 grains, but this is doubtful.

PAREIRÆ RADIX.

PAREIRA ROOT

The dried Root of *Chondrodendron tomentosum*, Ruiz and Pavon

Under the title of **Cissampelos**, the dried Root of *Cissampelos Pareira*, L., is official in the *Ind* and *Col* Add for India and the Eastern Colonies, also **Decoctum Cissampeli** (1 in 8), dose $\frac{1}{2}$ to 2 fl oz = 14.2 to 56.8 cc, and **Extractum Cissampeli Fluidum** (1 in 1), dose 30 to 120 minims = 1.8 to 7.1 cc.

Imported from Rio Janeiro in South Brazil. A spurious Pareira has lately been imported from Bahia in North Brazil, much inferior in alkaloid and extractive. The most marked chemical difference between the two is in the Petroleum Ether Extractive. In the genuine drug this amounts to over 8 p c, and in the spurious to about 0.3 p c.—*P J* (3) xxii 703, 771.

A good deal of the stem, which closely resembles the root, is also imported, and is said to be much less efficacious. Several drugs have been sold at different times as Pareira Brava.

Medicinal Properties—Astringent and mild diuretic in catarrhal affections of the genito-urinary tract, such as gonorrhœa and cystitis.

Official Preparation—**Extractum Pareiræ Liquidum**

Foreign Pharmacopœias—Official in Mex and Port (Butira), US Not in the others.

Descriptive Notes—Although the Root only is official in the *BP*, the stem usually comes with it into commerce, often in the proportion of 3 parts to 1 of root. The root is of a black colour externally, longitudinally furrowed, and marked with transverse ridges. The bark is thin. Internally the root is brownish or yellowish grey, with several more or less concentric zones having crenated edges, the porous woody wedges being separated by broad medullary rays, it has a waxy surface when cut. It has a bitter taste but no odour. The root is officially limited to pieces $\frac{1}{2}$ to 2 or more in (2 to 5 cm) in diameter. The stem is similar internally, but externally is of a pale greyish colour, with numerous warty, round lenticels. According to Moss the stem contains only three-fourths of

the amount of active principle yielded by the root. The true Pareira comes into commerce with considerable irregularity, but its place is taken by inert or false roots all of which have narrower and more numerous zones, which have not, in any case, crenated edges.

Tests—Pareira Root leaves from 3 to 4 p c of ash when ignited with free access of air, and the latter figure should not be exceeded. It has been stated [*P J* (3) xxii 703, 771] to contain about 8 p c of fatty matter extractable by Petroleum Ether, but genuine Pareira root and Bahia root obtained from an authoritative source did not yield that amount of Petroleum Ether extract. Samples of Pareira root examined in the author's laboratory yielded on an average 0.5 p c w/w of Petroleum Ether extract.

Preparation.

EXTRACTUM PAREIRÆ LIQUIDUM. LIQUID EXTRACT OF PAREIRA

Pareira Root exhausted with boiling Distilled Water, the liquid evaporated until it contains 33½ p c by weight of solid extract, then mixed with one-third of its volume of Alcohol (90 p c).

B P 1885 prepared the fluid extract from a solid extract.

Dose.—½ to 2 fl drms = 1.8 to 7.1 cc

Incompatibles—Ferric salts, Lead salts, Tincture of Iodine.

Foreign Pharmacopœias—Official in U S, 1 in 1 with Glycerin. Not in the others.

Tests.—Liquid Extract of Pareira has a sp gr of 1.025 to 1.048, it contains from 12 to 22 p c w/v of total solids and about 22 p c w/v of Absolute Alcohol.

PEPSINUM.

PEPSIN

FR, PEPSINE, GER, PEPSIN, ITAL, PEPSINA, SPAN, PEPSINA MEDICINAL

A proteolytic ferment or enzyme obtained from the mucous lining or the glandular layer of the fresh stomach of the healthy pig,* sheep, or calf. The *USP* admits only the fresh stomach of the hog.

A fine white or yellowish-white amorphous powder, or thin pale yellow or yellowish translucent grains or scales, without any offensive odour, and having a slightly saline taste, followed by a regurgitation of bitterness. It should be kept in well-closed glass bottles, as it slowly absorbs moisture when exposed to the air.

B P requires it to dissolve 2500 times and *USP* not less than 3000 times its own weight of freshly coagulated and disintegrated egg albumen.

Solubility.—Soluble about 1 in 100 of Water, more soluble in Water acidulated with Hydrochloric Acid. Insoluble in Alcohol (90 p c).

B P states that Pepsin is soluble 1 in 100 of Alcohol (90 p c), but this can only apply to the 500-test Pepsin of *B P* '85, because it consists principally of Sugar of Milk and not Pepsin. It is not true of Pepsin, *B P* '98.

Medicinal Properties—A digestive adjuvant, preferably given with dilute Hydrochloric Acid, used in chronic dyspepsia with deficiency of gastric juice, and in irritability of stomach associated with vomiting and gastralgia. It does not aid the digestion of carbohydrates and fats. It ought to be taken immediately after meals.

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Prescribing Notes—Given in powders, or in pills with 'Dispensing Syrup,' also in cachets, capsules, and compressed tablets

Official Preparation—Glycerinum Pepsini

Not Official—Elixir Pepsini, Elixir Pepsini et Bismuthi, Elixir Pepsini et Bismuthi Compositum, Elixir Pepsini et Bismuthi cum Ferro, Elixir Pepsini et Bismuthi cum Podophyllino, Elixir Pepsini et Bismuthi cum Strychnina, Elixir Pepsini et Bismuthi et Strychnina cum Ferro, Elixir Pepsini et Quinina, Elixir Pepsini et Quinina cum Ferro, Glycerole of Pepsin, Glycerinum Pepsini Fortius, Liquor Pepticus, Mistura Pepsini Composita, Mistura Pepsini cum Extracto Malti, Pepsinum Saccharatum, Vinum Pepsini

The usual solvent for making fluid preparations of Pepsin is a weak Alcohol acidulated with Hydrochloric Acid, to which Glycerin is added.

Alcoholic media are stated to be unsuitable vehicles for pharmaceutical preparations of Pepsin, as even dilute Alcohol in time destroys its activity—*L* '02, 1 687, *P J* '02, 1 294

Pepsin is one of the soluble ferments or enzymes of the gastric juice. It dissolves natural proteids, albumens, and fibrin, and converts them into syntonin and subsequently into albumose and Peptone. It is a conversion of the less soluble proteids into those that are more so, Peptone being the most soluble and diffusible of the proteids. Pepsin has no action on starch.

It acts only in acid solution, 0.2 p.c. of Hydrochloric Acid being the most favourable.

The action of Pepsin will continue almost indefinitely if the products of its action are removed by dialysis, or if the concentration of the products is reduced by acidified Water.

The gastric juice also contains another enzyme, 'rennin,' which curdles milk. The curd is formed in acid or neutral solutions in the presence of Calcium Phosphate. The casein is split up into a soluble and an insoluble proteid, the latter of which entangles the fat and forms a curd.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap (Pepsinum Saccharatum), Mex, Norw, Port, Russ, Span, Swed, Swiss and U.S.

Tests—Pepsin is required by the *B.P.* to dissolve 2500 times its weight of coagulated egg albumen and the following test is adopted for ascertaining that the sample possesses this degree of activity. A weighed quantity of 12.5 grammes of firm coagulated egg albumen prepared by boiling fresh eggs in Water for a quarter of an hour, chilling them in cold Water, separating and washing the whites free from pellicle or yolk, drying with a clean cloth and rubbing the coagulated Albumen through a sieve containing 12 meshes to a cm., is mixed with 125 c.c. of a 0.2 p.c. w/w Hydrochloric Acid Solution. A weighed quantity of 5 mg. of Pepsin is added and the whole digested for 6 hours at a temperature of 40.5° C (105° F), the mixture being shaken frequently. The coagulated egg albumen is officially required to dissolve to an almost clear solution leaving only a few small flakes. Care should be taken not to expose the coagulated white of egg to too long a contact with the atmosphere before starting the determination, as the product should be used before it has

lost moisture The above process has been subjected to severe and well-merited criticism, it has been objected that the quantity of 5 mg is an absurdly small one for the test, and that the *USP* method of employing an aliquot portion of a solution of Pepsin of a known strength is preferable The *USP* now allows the digestion to proceed for 2½ hours instead of for 6 hours, as in the 1891 edition, and give the exact directions as to the number of times and the method by which the liquid is to be agitated, but neither the *BP* nor the *USP* makes any allowance for the solvent action of the acid on the albumen Allen shows that by only requiring the Pepsin to dissolve the albumen no distinction is drawn between its conversion into Syntonin and true peptonisation The real digestive power of a Pepsin is measured by the amount of Peptone which it produces in a given time under certain conditions The conditions of the *USP* method of experiment afford a determination of the solvent action of the Pepsin on the albumen, although the *USP* description states that when assayed by such process it shall be capable of digesting the albumen Allen has worked out a process, which he claims to be an original one for the determination of the digestive power of a Pepsin, whereby the actual amount of Peptone or of mixed Peptones and Albumose produced by digestion is ascertained The process may be roughly outlined as follows —A weighed quantity of about 1 gramme of scale egg-albumen is powdered and treated with 20 c.c of warm Water, and when dissolved is heated in a water-bath to coagulate the albumen and cooled to a temperature not exceeding 40° C (104° F), 0.1 of a gramme of a sample of Pepsin to be tested is added, followed by 25 c.c of Tenth-normal Volumetric Hydrochloric Acid Solution, the mixture is warmed to 40° C (104° F) and maintained at this temperature for 3 hours A volume of Tenth-normal Volumetric Sodium Carbonate Solution exactly equivalent to the Tenth-normal Volumetric Acid Solution previously used is then added and the liquid heated on a water-bath to 90° C (194° F) for 10 minutes, it is cooled, diluted with Water to 100 c.c and passed through a fine filter, the precipitate contains Syntonin and any unaltered albumen, the filtrate containing the Albumose and Peptones A measured quantity of 50 c.c is saturated in the cold with Zinc Sulphate (about 60 grammes being required for 50 c.c) and the mixture is allowed to stand for half an hour with intervals of occasional stirring and filtered, the precipitate washed with cold saturated Zinc Sulphate Solution, diluted with Water to 150 c.c, acidulated with Hydrochloric Acid and treated with Bromine Water, the precipitate is filtered and the Nitrogen determined by Kjeldahl's test, allowance being made for the Nitrogen contained in the Pepsin employed The method is described in the *Analyst* '97, 258

The *USP* Pepsin is required to 'digest' not less than 3000 times its own weight of freshly coagulated and disintegrated egg albumen, and a method of determination of which the following are the essential features is employed —The coagulated albumen is prepared on somewhat similar lines to the *BP*, the coagulated albumen when separated from the pellicle and yolk being

rubbed through a clean No 40 sieve, the first portions passing through being rejected and a weighed quantity of 10 grammes of the succeeding portion transferred to a wide mouthed bottle of 100 cc capacity. A measured quantity of 9 cc of Diluted Hydrochloric Acid is mixed with 201 cc of Water, and in 150 cc of this acid liquid a weighed quantity of 1 dg of Pepsin is dissolved. A measured quantity of 20 cc of the Diluted Hydrochloric Acid liquid is added to the 10 grammes of albumen in the bottle and the albumen completely disintegrated by rubbing with a rubber-tipped glass rod, the rod being rinsed with a further quantity of 15 cc of diluted acid liquid, and finally a measured quantity of 5 cc of the solution of Pepsin is added. After the bottle has been securely corked, it is inverted three times and maintained at a temperature of 52° C (125 6° F) for 2½ hours, the bottle being inverted once in 10 minutes. At the end of this time the source of heat is removed, 50 cc of cold Water added, the mixture transferred to a narrow graduated cylinder and allowed to remain at rest for half an hour. The *USP* requires that the precipitate of undissolved albumen should not measure more than 1 cc. The relative proteolytic power of a stronger or weaker Pepsin may be determined by a series of experiments to ascertain the exact quantity of a Pepsin solution required on the lines prescribed above, to digest the 10 grammes of coagulated and disintegrated egg albumen, the quantity in cc of Pepsin Solution required divided into 15,000 gives the number of parts of egg albumen digested by one part of Pepsin.

The *PG* test is made on the following lines — A weighed quantity of 10 grammes of disintegrated egg albumen, prepared from an egg which has been boiled for 10 minutes, and after separation of the yolk, the white has been reduced to a state of coarse powder by rubbing through a sieve, is mixed with 100 cc of warm Water of a temperature of 50° C (122° F), and a measured quantity of 0.5 cc. of Hydrochloric Acid added, and finally a weighed quantity of 0.1 of a gramme of Pepsin, the mixture is allowed to stand for 1 hour at 45° C (113° F) with repeated intervals of shaking. The *PG* requires that, with the exception of a few yellowish-white particles, the albumen shall be completely dissolved.

A method based on the lines of the *USP* has been suggested (*PJ* '04, u 376) the solution of Pepsin was prepared by triturating 25 cg of Pepsin, 1 gramme of Sodium Chloride, and adding acidulated Water very carefully at first, mixing well and transferring to 1000 cc flask, the containing vessels being rinsed out with acidulated Water and made up to 1000 cc, the solution is allowed to stand for 24 hours and shaken at intervals. A measured quantity of 20 cc is placed in a stoppered bottle of about 250 cc capacity, and 12 grammes of coagulated egg albumen, which has been previously thoroughly reduced to uniform granules by trituration in a small mortar with 50 cc of acidulated Water, is added, the last traces of albumen being transferred to the flask with a further quantity of 50 cc of acidulated Water, the mixture is digested at a temperature of 45° C (113° F) for 6 hours, with intervals of vigorous shaking for 15 minutes.

Preparation

GLYCERINUM PEP SINI.—GLYCERIN OF PEP SIN

Hydrochloric Acid, 110 minims, Glycerin, 12 fl oz, Distilled Water, 6 fl oz, Pepsin, 800 grains, macerate for a week, filter and make up with Distilled Water to 20 fl oz (1 in 11)

Dose.—1 to 2 fl drms = 3 6 to 7 1 c c, corresponding to 5 to 10 grains = 0 32 to 0 65 grammes of Pepsin

The Pepsin should be dissolved in the Water, the Glycerin added in 3 or 4 portions, with agitation, then the Acid, and finally made up to volume with Water, and filtered.—*P J '04, 1 84*

Not Official

PEPSINUM SACCHARATUM—Pepsin, 1, Sugar of Milk, recently dried and in No 30 powder, 9—*U S P 1890 and Jap*

ELIXIR DE PEP SINE—Pepsin, 2, Distilled Water, 28, Vin de Lunel, 50, Glycerin, 20—*Fr*

ELIXIR PEP SINI—Pepsin, 5, Alcohol, 15, Distilled Water, 45, Aromatic Elixir, *q s* to produce 100—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI** *Syn* Bismuth and Pepsine Mixture—Stronger Glycerin of Pepsin, 12 50, Bismuth and Ammonium Citrate, 3 50, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100 Mix the Glycerin of Pepsin with 10 of the Simple Elixir, and neutralise the mixture carefully with a weak solution of Ammonia Dissolve the Bismuth and Ammonium Citrate in 50 of the Simple Elixir, adding solution if acid by neutralising with Ammonia Finally mix the two solutions, add the Alcohol, make up the required volume with Simple Elixir, and filter **Dose**— $\frac{1}{2}$ to 1 fl drms = 1 8 to 3 6 c c—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI COMPOSITUM**—Stronger Glycerin of Pepsin, 12 50, Bismuth and Ammonium Citrate, 3 50, Morphine Acetate, 0 10, Diluted Acetic Acid, 0 20, Tincture of Nux Vomica, 2, Diluted Hydrocyanic Acid, 2, Alcohol (60 p c), 5, Solution of Cochineal, 2 5, Simple Elixir, *q s* to produce 100 Mix the Glycerin of Pepsin with 10 of the Simple Elixir, and neutralise the mixture carefully with a weak solution of Ammonia Dissolve the Bismuth and Ammonium Citrate in 50 of Simple Elixir, adding solution of acid by neutralising with Ammonia Next mix the Acetic Acid, Alcohol, and 5 of the Simple Elixir, and dissolve the Morphine Acetate in the mixture Mix the three solutions, add the tincture of Nux Vomica, then the Hydrocyanic Acid, and sufficient Simple Elixir to make up the required volume Finally colour with the solution of Cochineal, and filter **Dose**— $\frac{1}{2}$ to 1 fl drms = 1 8 to 3 6 c c—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI CUM FERRO**—Stronger Glycerin of Pepsin, 12 50, Bismuth and Ammonium Citrate, 3 50, Iron and Ammonium Citrate, 3 50, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI CUM PODOPHYLLINO**—Stronger Glycerin of Pepsin, 12 50, Bismuth and Ammonium Citrate, 3 50, Podophyllum Resin, 0 25, Aromatic Spirit of Ammonia, 5, Solution of Cochineal, *q s*, Simple Elixir, *q s* to produce 100—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI CUM STRYCHNINA**—Stronger Glycerin of Pepsin, 12 50, Bismuth and Ammonium Citrate, 3 50, Solution of Strychnine Hydrochloride, 2 50, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100—*B P C*

* **ELIXIR PEP SINI ET BISMUTHI ET STRYCHNINÆ CUM FERRO**—Stronger Glycerin of Pepsin, 12 50 Bismuth and Ammonium Citrate, 3 50, Solution of Strychnine Hydrochloride, 2 50, Iron and Ammonium Citrate, 2, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100—*B P C*

* These formulas closely resemble those previously published in *Armour's Formulary* (9th edit)

* **ELIXIR PEPSINI ET QUININÆ** —Stronger Glycerin of Pepsin, 12 50, Quinine Acid Hydrochloride, 1, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100 —*B P C*

* **ELIXIR PEPSINI ET QUININÆ CUM FERRO** —Stronger Glycerin of Pepsin, 12 50, Iron and Quinine Citrate, 3 50, Alcohol (60 p c), 5, Simple Elixir, *q s* to produce 100 —*B P C*

ELIXIR PEPSIN ET EUONYMIN See p 500

GLYCEROLE OF PEPSIN —Pepsin, 2 oz, Diluted Hydrochloric Acid, 1 fl oz, Glycerin, 10 fl oz, Simple Elixir, 1 fl oz Distilled Water, *q s* to produce 20 fl oz —*P J F*

GLYCERINUM PEPSINI FORTIUS *Syn* Glycerol of Pepsin —Pepsin, 15, Diluted Hydrochloric Acid, 5, Glycerin, 50, Simple Elixir, 5, Distilled Water, *q s* to produce 100 —*B P C*

* **LIQUOR PEPTICUS** —Stronger Glycerin of Pepsin, 12 50, Diluted Hydrochloric Acid, 2 50, Alcohol, 10, Glycerin, 2 50, Distilled Water, *q s* to produce 100 —*B P C*

* **MISTURA PEPSINI COMPOSITA** —Stronger Glycerin of Pepsin, 5, Solution of Strychnine Hydrochloride, 1 25, Diluted Nitro Hydrochloric Acid, 3, Glycerin, 10, Tincture of Cudbear, 5, Distilled Water, *q s* to make 100 —*B P C*

* **MISTURA PEPSINI CUM EXTRACTO MALTI** *Syn* Essence of Pepsin and Malt —Stronger Glycerin of Pepsin, 5, Extract of Malt, by weight, 30, Alcohol (60 p c), *q s* to produce 100 —*B P C*, altered in *B P C Supp*

* **Elixir Simplex** —Tincture of Orange, 7 50, Syrup, 40, Distilled Water, *q s* to produce 100 Mix the Tincture with the Syrup, add sufficient Distilled Water to make up the required volume, and filter through kaolin —*B P C*

* **Tinctura Persionis** Tincture of Cudbear —Cudbear, in fine powder, 12 50, Alcohol, 35, Distilled Water, *q s* to produce 100 —*B P C*

VINUM PEPSINI Pepsin Wine —Pepsin, 320 grains, Hydrochloric Acid, 2 fl drn, Glycerin, 1 fl oz, Sherry, *q s* to 20 fl oz —*B P C Formulary* 1901, now incorporated in the *B P C*, with 3½ p c of Pepsin and using Detannated Sherry, as follows —

Pepsin, 3 50, Hydrochloric Acid, 1 25, Glycerin, 5, Detannated Sherry, *q s* to produce 100

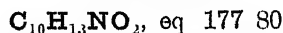
Official in Austr, Belg, Dutch, Ger and Russ, about 1 in 40, Jap about 1 in 20, Mex, 1 in 30, Span, 1 in 20

Ingluvin —An amorphous powder, prepared from the gizzard of the domestic fowl Introduced as a substitute for Pepsin A stomachic tonic for the relief of indigestion, flatulence and dyspepsia, and of special use in the vomiting of pregnancy

Dose —5 to 10 grains = 0 32 to 0 65 grammo

PHENACETINUM.

PHENACETIN



FR, ONIHYLIARA ACETIANIDE, GER, PHLNACETIN, ITAL, FENACMINA, SPAN, FENACETINA

A white, odourless, almost tasteless, crystalline powder, or white glistening crystalline scales It is produced by the action of Glacial Acetic Acid upon Para-phenetidin

* These formulas closely resemble those previously published in *Armon's Formulary* (9th edit)

Phenacetin is described in the *USP* under the title of Acetphenetidin and is stated to be a Phenol derivative, the product of the acetylation of Para-amidophenetol

Solubility.—1 in 1700 of Water, 1 in 50 of boiling Water, 1 in 21 of Alcohol (90 p c), 1 in 100 of Alcohol (60 p c)

Medicinal Properties.—Analgesic, antipyretic and nervine sedative. It does not produce nausea, and it depresses the heart very little, when used judiciously. It is an efficient synthetic analgesic for the relief of neuralgic, rheumatic, locomotor ataxial and other pains, and is the safest of the synthetic antipyretics, being the most free from toxic effects.

As the result of an inquiry as to the ill-effects of Phenacetin, by a Committee of the British Medical Association, it is stated that it appears to have a notable freedom from injurious action, and has great value, especially as an analgesic. Some observers recommend a commencing dose of 5 grains or less, others using doses of 8 to 10 grains.—*B M J* '94, i 89

Two cases of temporary rash caused by Phenacetin without any other untoward result.—*L* '95, i 91, *C D* '95, i 797

Palpitation caused in an adult male by taking 8 15-grain powders in 3½ hours.—*Pr* li 241, palpitation caused in a female aged thirty-two by taking 5 to 8 cachets of 10 grains each in the 24 hours.—*Pr* liii 444

Phenacetin and Antipyrine are the most trustworthy and valuable of this class of pain-relieving remedies, and if used with due care and judgment, ill-effects following the use of either are exceedingly rare, the principal precaution being to commence with a small dose, of Phenacetin 5 grains and of Antipyrine not more than 10 grains.—*Scot Med and Surg Jour* '98, ii 486

Recommended in influenza to relieve the headache and reduce temperature.—*B M J* '91, i 1282, '91, ii 190, '94, ii 1045

Dose.—5 to 10 grains = 0.32 to 0.65 gramme

Ph Ger maximum single dose, 1.0 gramme, maximum daily dose, 3.0 grammes

Prescribing Notes.—It is given in cachets, or suspended in Water with Compound Powder of Tragacanth, in migraine it is usually given with Caffeine in effervescent granules

Not Official.—Phenacetinum cum Caffeina Effervescent, Amygdophenin, Eupyrine, Kryofin, Lactophenin, Malakin, Para-phenetidin and Citrates, Phenosal, Triphenin, Phenocoll Hydrochloridum and Salocoll

Foreign Pharmacopœias.—Official in Belg, Dan, Dutch, Ger, Jap, Norw, Russ, Swed and Swiss (Phenacetinum), Austr and U.S. (Acetphenetidinam), Fr (Oxethylpara-Acétanilide), Ital, Mex and Span (Genacetina)

Tests.—Commercial Phenacetin melts at 133–79° C (272–82° F), the dried product at 134–26° C (273–67° F), and the purified product at 134–89° C (274–80° F). The *BP* m.p. is 135° C (275° F). The *USP* and *PG* m.p. is 134° to 135° C (273–2° to 275° F). It dissolves in Sulphuric Acid without change of colour. When 0.1 of a gramme is boiled for from half a minute to 1 minute with 1 or 2 c.c. of Hydrochloric Acid it yields a fluid which, when diluted with 10 times its volume of Water, yields on the addition of 3 or 4 drops of Chromic Acid Solution a deep red coloration. The *USP* states that it is coloured yellow by Nitric Acid, the colour persisting when heated.

The more generally occurring impurities are Acetanilide, uncon-

verted Para-phenetidin, and mineral matter The *BP* gives only one test for the detection of Acetanilide, requiring that a cold saturated aqueous solution of the sample should not be rendered turbid by the addition of Bromine Solution This test is also given in the *PG* and *USP* The *USP* gives three separate tests for Acetanilide the Potassium Hydroxide test, the Sodium Hydroxide and Chlorinated Soda test, and the Bromine test described in small type below The Iso-nitrite test is not included, this latter test, when carried out according to the modification described under Acetanilide, is capable of detecting readily an addition of 2 pc of the latter substance An admixture of Acetanilide also affects the mp, pure Phenacetin and pure Acetanilide did not begin to fuse at any temperature approaching 92° C (197.6° F), whereas mixtures containing from 1 to 95 pc of Acetanilide all commenced to fuse at this temperature Unconverted Para-phenetidin, if present, may be detected by the reddish tint developed on the addition of Volumetric Iodine Solution The *BP* and the *USP* require that a mixture of 0.3 of a gramme of Phenacetin with 1 cc of Alcohol when diluted with 3 times its volume of Water should not acquire a red coloration on boiling with 1 drop of Volumetric Iodine Solution It may also be detected by the dark red colour produced on the addition of Ferric Chloride T.S. to the saturated aqueous solution of the sample Phenacetin when heated with free access of air should leave no weighable residue

Bromine—The solution obtained when 0.1 gramme is boiled with 10 cc of Water, cooled and filtered, should not be rendered turbid by the addition of a slight excess of Bromine Water, *PG* and *USP*

Potassium Hydroxide—0.1 gramme, heated with 5 cc of a solution of Potassium Hydroxide (1 in 4), should not give off a perceptible odour of Aniline, *USP*

Sodium Hydroxide and Chlorinated Soda—Boil 0.1 gramme for 1 minute with 8 cc of a solution of Sodium Hydroxide (1 in 2), cool, and agitate the solution with 5 cc of solution of Chlorinated Soda A clear yellow liquid should be obtained, and not a purplish red or brownish red cloudy liquid or precipitate, *USP*

Not Official

PHENACETINUM CUM CAFFEINA EFFERVESCENS—Sodium Bicarbonate, 46, Tartaric Acid, 24, Citric Acid, 16, Refined Sugar, 16½, Phenacetin, 5, Caffeine Citrate, 2½, make into granules of a suitable size—*BPC* (about 5 in 100)
Formulary 1901

This has been incorporated in the *BPC*, but the Caffeine is reduced to 1½ and the Sugar to 16, *BPC Supp* has altered the Citric Acid to 18, and the Sugar to 15½

Dose—60 to 120 grains = 4 to 8 grammes

AMYGDOPHENIN (Para-phenetidin Amygdalate)—A greyish white, voluminous, crystalline powder, very sparingly soluble in Water Anti-rheumatic and antineuralgic, but of little value as an antipyretic—*PJ* '93, 189, 162, *BMJE* '95, 11 99

Dose—8 to 15 grains = 0.52 to 1 gramme

CITROPHEN (Para-phenetidin Citrate)—A white powder, with an acid reaction, soluble 1 in 165 of Water, 1 in 300 of Alcohol (90 pc), insoluble in Ether and in Chloroform Antipyretic and analgesic, sometimes causing considerable sweating

Useful in rheumatism of the joints and muscles, in the severe headache of influenza, and in acute tonsillitis —*B M J E* '99, 11 52

Dose — $\frac{7}{8}$ to 15 grains = 0.5 to 1 gramme

Tests —Citrophen possesses a m.p. of about 181° C (357.8° F). A small quantity, when boiled with Hydrochloric Acid, cooled, and the solution diluted with Water, yields a liquid which acquires a deep reddish colour on the addition of Chromic Acid Solution. When ignited with free access of air it should leave no weighable residue.

Citrophen is dibasic, and Apolysin is monobasic, Para-phenetidin Citrate

Apolysin forms yellowish-white crystals, or a crystalline powder, with an acid reaction, readily soluble in Water. Has been used as an antipyretic and analgesic.

EUPYRINE (Para-phenetidin-vanillin-ethyl Carbonate) —Pale, greenish-yellow crystals, insoluble in Water, readily soluble in Alcohol (90 p.c.), in Ether and in Chloroform. Introduced as an innocuous antipyretic —*P J* '01, 11 312, *C D* '01, 1 36

Dose —15 to 20 grains = 1 to 1.3 grammes

KRYOFIN (Para-phenetidin Methylglycollate) —White, odourless, tasteless crystals, sparingly soluble in cold Water. Antipyretic and analgesic. Useful in neuralgia. Severe sweating sometimes follows its use —*B M J E* '97, 1 83, '97, 11 88, *L* '97, 11 728, *P J* '97, 11 5

Dose —8 to 15 grains = 0.52 to 1 gramme

LACTOPHENIN (Para-phenetidin Lactate) —A white, inodorous, bitter, crystalline powder, sparingly soluble in Water.

Medicinal Properties —Antipyretic, analgesic and hypnotic. Used in migraine, erysipelas, nervous headache and the neuralgia of influenza.

Dose —5 to 15 grains = 0.32 to 1 gramme

Tests —Lactophenin melts at about 118° C (244.4° F). 0.1 gramme boiled with 1 c.c. of Hydrochloric Acid, the solution cooled and diluted with 10 c.c. of Water and filtered yields, on the addition of 1 or 2 drops of Chromic Acid Solution, a deep red coloration. It dissolves in Sulphuric Acid without change of colour. When ignited with free access of air it should leave no weighable residue.

MALAKIN (Para-phenetidin Salicylate) —Occurs in pale yellow, silky needles or a yellow, crystalline powder, insoluble in Water and strong Alcohol.

At pyretic analgesic and antirheumatic. Used in acute rheumatism, the fever of phthisis, migraine and neuralgia —*M P* '94, 1 268, *J* '94, 1 92, '94, 1 84, '94, 11 88, *T G* '95, 325, *Pr* lin 45, *Y B P* '95, 1 89, in every way inferior to Phenacetin and Antipyrine —*B M J* '95, 1 5

Dose —10 to 20 grains = 0.65 to 1.3 grammes

PARA-PHENETIDIN CAMPHORATE —A white, crystalline powder, insoluble in Water, soluble in Alcohol (90 p.c.). Introduced as an antipyretic.

PHENOSAL (Para-phenetidin Aceto-salicylate) —A white, odourless, crystalline powder, sparingly soluble in Water, in Alcohol (90 p.c.) and in Ether. It is stated to possess antipyretic and antineuralgic properties —*P J* '99, 11 62.

Dose —5 to 10 grains = 0.32 to 0.65 gramme

Triphenin —A derivative of Para-phenetidin and Propionic Acid, dose, 8 to 15 grains. **Phesin**, a sulfo-derivative of Para-phenetidin, and **Pyranthin**, a derivative of Para-phenetidin and Succinic Acid, dose 5 to 10 grains, have been recommended as antipyretics. **Chinaphenin** (Para-phenetidin-quinine-ethyl Carbonate), dose 5 to 15 grains, and **Para-phenetidin Agarate** have been introduced as antipyretics.

PHENOCOLL HYDROCHLORIDUM —A white, crystalline powder soluble 1 in 16 of Water, sparingly soluble in Alcohol (90 p.c.). Obtained by the action of Glycocoll on Phenetidin.

Medicinal Properties—Antipyretic, yielding good results in rheumatic fever—*L* '91, 1 1060, '92, 11 438. As a substitute for Quinine in malaria, *B M J E* '93, 11 104, *T G* '93, 334, 618, in acute rheumatism, typhoid, malaria, and as an intestinal antiseptic, *B M J E* '94, 1 79, '96, 11 83, *L* '97, 1 1227, *P J* '96, 1 178, used in 400 cases of influenza during an epidemic, and found to be a specific—*P J* '99, 11 216

Dose—5 to 10 grains = 0.32 to 0.65 gramme

Salocoll (Phenocoll Salicylate), recommended in rheumatism in doses of 15 to 30 grains = 1 to 2 grammes. It is not so soluble in Water as the Hydrochloride

PHENAZONUM.

PHENAZONE

$C_{11}H_{12}N_2O$, eq 186.77

FR, ANTIPYRIN, GER, PHENYLDIMETHYLPYRAZOLON, ITAL, ANTIPIRINA, SPAN, ANTIPIRINA

Colourless, odourless, crystalline scales, or as a white, neutral, odourless powder, possessing a somewhat bitter taste

B P states that 'Phenazone is commonly known as Antipyrine,' but it is not very clear from this note whether it is intended that Phenazone should be used when Antipyrine is ordered, or that Antipyrine should be used when Phenazone is ordered, or whether it is an incidental note having no meaning

It appears in the *U S P* under the title of Antipyrina, and is stated to be obtained by the condensation of Phenylhydrazine with Acetoacetic Ether and subsequent methylation

Solubility—1 in $1\frac{1}{4}$ of Water, 3 in 4 of Alcohol (90 p c), about 5 in 6 of Chloroform, 1 in 40 of Ether

Medicinal Properties—Antipyretic and analgesic, nervine sedative. It will reduce temperature in all forms of febrile disease, but in weak subjects its depressant effect should be borne in mind

As an analgesic it is used with great success in neuralgia, migraine, gout, rheumatism, locomotor ataxia and other painful affections, and is frequently given with Sodium Salicylate and Caffeine

It is a good uterine sedative, it also relieves sea-sickness

As a pain-relieving remedy Phenacetin is preferred by some, as it is less likely to produce toxic effects

Of great value as a sedative in some of the nervous disturbances of childhood—*P* '07, 1 540

10 p c solution locally in epistaxis—*M A* '94, 253, *L* '93, 11 453. As a styptic and antiseptic—*B M J E* '95, 1 28, *L* '95, 1 1453. In Tannic Acid Solution as a styptic—*B M J E* '95, 11 90. One of the most pleasant and rapid remedies for influenza—*Pr* liv 383. Discussion on its benefits and risks as an analgesic—*B M J* '98, 11 1054, it is contra indicated in cardiac weakness and cases of extreme exhaustion—*T G* '89, 457

As the result of an inquiry as to the ill effects of Phenazone, by a Committee of the British Medical Association, it is stated that the commencing dose should not exceed 10 grains, and should not be repeated too frequently, there is a necessity for watching its action, but ill effects are not of the frequency or importance ascribed to them by a widespread impression. The large majority of observers agree in stating that they are of no importance whatever, and that,

with reasonable and judicious care, they limit in no way the general usefulness of the drug as a therapeutic agent.—*B M J*, '94, 1 88

Toxic symptoms following the administration of 10 grains dissolved in 1 oz of Water, recovery.—*B M J*, '99, 11 85.

Dose.—5 to 20 grains = 0.32 to 1.3 grammes—

Swiss, maximum single dose, 2.0 grammes, maximum daily dose, 6.0 grammes

Prescribing Notes—*Given in solution with Tincture of Orange and Spirit of Chloroform or Chloroform Water, or in powders, cachets, capsules, or in the form of effervescent granules*

Incompatibles—*Spiritus Ætheris Nitrosi*, Tannic Acid in aqueous solutions, *Extractum Cinchonæ Liquidum*, and other astringent decoctions and infusions. Chloral Hydrate is not incompatible with Phenazone in moderately dilute aqueous solution. Sodium Salicylate is not incompatible with Phenazone in aqueous solution, but forms an oily liquid if the *solids* be mixed and exposed to the air.—*P J* (3) x 861

The incompatibility of Antipyrine and *Spiritus Ætheris Nitrosi* may be overcome by prescribing them with Sodium Bicarbonate.—*A J P*, '94, 321, *C D*, '98, 1 857

Not Official—*Pulv Phenobrom Co*, *Phenazonum Effervescens*, *Acetopyrine*, *Feripyrin*, *Hypnal*, *Iodopyrin*, *Migramine*, *Pyramidon*, *Pyramidon Camphorates* and *Salicylate*, *Salpyrin*, *Tolpyrin*, *Tolysal* and *Tussol*

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Hung., Ital., Jap., Mex., Norw., Russ., Span., Swed., Swiss and U.S. (*Antipyrinum*), Ger. (*Pyrazolonum Phenylidimethylicum*)

Chloroform extracts Antipyrine from alkaline solution, but imperfectly from acid solution

Tests.—Commercial Phenazone melts at 109.12° C (228.42° F), dried Phenazone melts at 110.01° C (230.02° F), and purified Phenazone melts at 112.84° C (235.11° F). The *BP* states about 113° C (235.4° F), the *USP* and *PG* state that it melts at 113° C (235.4° F). The *Fr Codex* gives 114° C (237.2° F) as the m.p. 5 c.c. of an aqueous 1 p.c. solution of Phenazone when mixed with 5 c.c. of Nitric Acid, develops a yellow colour which, on warming, changes to crimson. 12 c.c. of a solution of similar strength, when mixed with 1 decigramme of Sodium Nitrite, yields an almost colourless fluid which, on the addition of 1 c.c. of Diluted Sulphuric Acid, assumes a deep green colour. In the place of Sodium Nitrite, a few drops of *Spiritus Ætheris Nitrosi* may be used, and will answer the same purpose. A very dilute solution affords, with Ferric Chloride T.S., a deep red coloration, the colour being destroyed by an excess of diluted Sulphuric Acid. The *USP* and *PG* recommend the use of 2 c.c. of a 1 in 1000 Phenazone solution and 1 drop of Ferric Chloride T.S. The *BP* states that the colour is nearly destroyed by excess of diluted Sulphuric Acid. The *USP* and *PG* state that it is changed to light yellow on the addition of 10 drops of Sulphuric Acid. The aqueous solution affords with Tannic Acid an abundant white precipitate, the *BP* states that a 5 p.c. aqueous solution affords with Mercuric Chloride T.S. a white precipitate, disappearing on boiling but reappearing on cooling. 2 c.c. of a 1 p.c. aqueous solution yields on the addition of 2 drops of Fuming Nitric Acid a green coloration, changing to red on boiling. The test is common to the *BP* and the *PG*, but is not in the *USP*. In an acidified aqueous solution it

yields a precipitate with Mayer's reagent, and also with Iodo potassium Iodide (Wagner's) Solution, in a similar way to an alkaloid. It may be extracted from alkaline solution by Chloroform, but is only imperfectly extracted from an acid solution. The *Ft. Codex* (1908) gives a method by which Antipyrine may be determined.—A weighed quantity of 0.5 of a gramme dissolved in 50 grammes of Water is mixed with 1.5 grammes of Sodium Acetate, and after having been brought to the boiling point is mixed with Iodo potassium Iodide Solution (4 grammes of Iodine and 2 grammes of Potassium Iodide dissolved in 10 grammes of Water and diluted with Water to 100 grammes), until the boiling liquid just commences to become coloured, when cooled the crystallised Iodine compound is separated, dried and weighed, 1 gramme represents 0.7 of a gramme of Antipyrine. It should be free from acids and alkaloids, as indicated by the behaviour of its aqueous solution towards Litmus paper. Its aqueous solution should not be affected by Hydrogen Sulphide, and it should be free from Acetanilide. Neither the *B.P.* nor the *P.G.* includes a specific test for Acetanilide, the *U.S.P.* employs the Iso nitrile test as described in the small type below. When ignited with free access of air it should leave no weighable residue.

Hydrogen Sulphide—The aqueous solution should not be affected by T.S. of Hydrogen Sulphide, *B.P.* and *P.G.*

Iso-nitrile Test—If 0.1 gramme be warmed with Sodium Hydroxide, Chloroform added and again warmed, the disagreeable odour of Phenyl isocyanide should not be developed, *U.S.P.*

Not Official

PULV. PHENOBROM. CO. (*Squre*)—A granular effervescent preparation, containing 5 grains of Phenazone, with Sodium Salicylate, Potassium Bromide, and Caffeine, in the tablespoonful dose.

PHENAZONUM EFFERVESCENS—A granular effervescent preparation, containing 8 grains of Phenazone in 100.—*B.P.C. Formulary* 1901.

Dose—60 to 120 grains = 4 to 8 grammes.

This has been incorporated in the *B.P.C.* under the title Antipyrina Effervescens, it also includes Antipyrina Effervescens cum Caffeina, containing, in addition, 1.5 grains in 100 of Caffeine Citrate.

The general composition is similar to that given under Phenacetin Effervescens.

ACETOPYRINE (Antipyrine Aceto salicylate)—A white, crystalline powder, sparingly soluble in Water, readily soluble in Alcohol (90 p.c.), possessing antipyretic and analgesic properties, recommended in rheumatism and neuralgia, and as a good antipyretic having no injurious action on the heart.—*M.P.* '04, 11 466.

Dose—5 to 10 grains = 0.32 to 0.65 gramme.

FERRIPYRIN—A compound of Antipyrine and Ferric Chloride, containing about 64 p.c. Antipyrine. Occurring as an orange red powder, soluble in Water. In 20 p.c. solution it has been found useful as a styptic. Useful in chlorosis and anæmia.—*B.M.J.* '95, 1 1382, *L.* '95, 1 1320, *B.M.J.E.* '95, 1 44, as analgesics, Ferripyrin, Tolpyrin and Pyramidon appear to be neither beneficial nor harmful, and are therefore of no therapeutic value for the relief of urgent pain.—*Scot. Med. and Surg. Jour.* '96, 11 442.

Dose—Usually 5 grains = 0.32 gramme.

HYPNAL—Is a crystalline compound of Antipyrine with Chloral Hydrate, readily soluble in Water, has been recommended as a hypnotic, used in simple

insomnia, delirium tremens and maniacal excitement—*Pr* 1 297, in the insomnia due to neuralgia or migraine, or the pyrexia of phthisis—*M P* '94, 1 267

Dose—10 to 20 grains = 0 65 to 1 3 grammes

It possesses the depressing action on the heart of both Antipyrine and Chloral Hydrate

IODOPYRIN—Colourless, glistening, prismatic needles, or as a white, crystalline powder, sparingly soluble in Water, soluble in Alcohol (90 p c) Antipyretic and antiseptic Has been given in puerperal fever, and has been found useful in acute articular rheumatism—*B M J E* '00, 1 12

Dose—5 to 15 grains = 0 32 to 1 gramme

Bromopyrin has also been given in the same doses as an antipyretic

MIGRAININE—A registered name for a double Citrate of Caffeine and Antipyrine A white, odourless powder, soluble in Water Has been found useful in migraine and in neuralgia—*C D* '95, 1 3, *P J* '97, 1 18

Dose— $7\frac{1}{2}$ to 15 grains = 0 5 to 1 gramme

Official in Austr and Swiss, Antipyrinum Caffeina-citricum

PYRAMIDON (Dimethyl-amido antipyrin)—A yellowish-white, tasteless, crystalline powder, readily soluble in Water and in Alcohol (90 p c) It is official in *Fr Codex* (1908) under the title of **Diméthylamino-antipyrine**, $C_{13}H_{17}N_3O$

Dose—As an analgesic, 10 grains = 0 65 gramme, as an antipyretic, 3 to 5 grains = 0 2 to 0 32 gramme—*B M J E* '97, 1 7, 84, '00, 1 56

5 grains several times daily in asthma, especially when of reflex origin—*P J* '03, 1 340

In typhoid fever 5 to 6-grain doses regularly night and day, every 2 hours, until temperature keeps down without it, in severe cases 6 grains, in mild 3 grains—*B M J E* '03, 1 23

4 grains repeated in half an hour caused marked reduction in temperature in typhoid fever—*B M J E* '03, 1 79

In 10 to 15-grain doses 3 or 4 times daily is of great value (*B M J E* '05, 1 72, '07, 1 3) in the treatment of typhoid when there is no contra-indication

Tests—Pyramidon melts at about 108° C (226 4° F), and this m p is given in *Fr Codex* The aqueous solution affords, on the addition of Ferric Chloride T S, a bluish-violet coloration, and on the addition of Sodium Nitrite and diluted Sulphuric Acid, and also on the addition of Fuming Nitric Acid, it affords a fluorescent bluish-violet coloration *Fr Codex* requires that a weighed quantity of 0 5 gramme dissolved in 50 c c of Water should, if pure, neutralise 21 75 c c of Normal Volumetric Sulphuric Acid Solution, whilst a similar weight of the official salt should neutralise more than 20 c c, Methyl Orange Solution being used as an indicator of neutrality The solution intended is evidently Dec i-normal When heated with free access of air it should leave no weighable residue

Pyramidon Mono- and Bi-Camphorate and -Salicylate.—The two former are antipyretic and anhidrotics in doses of 5 to 10 grains, the latter has been found useful in subacute and chronic rheumatism, also in doses of 5 to 10 grains = 0 32 to 0 65 gramme All occur as white crystalline powders The urine of patients taking Pyramidon is stated (*P J* '05, 1 270) to contain a new acid—Rubazonic Acid—which forms crystalline needles insoluble in Water

SALIPYRIN Antipyrine Salicylate $C_7H_7N_3O$ $C_7H_5O_3$, eq 328 78—Hexagonal crystals, or a white, crystalline, odourless powder, soluble 1 in 240 of Water, soluble 1 in 4 of Alcohol (90 p c)

In uterine hæmorrhage, *B M J E* '93, 1 82, *L* '95, 1 1005, *P J* '95, 1 363, a specific for influenza, *Y B T* '95, 454, *B M J E* '93, 1 103, in pelvioid rheumatism, *B M J E* '97, 1 44, analgesic in painful rheumatic conditions, *B M J* '98, 1 1055

Dose—10 to 30 grains = 0 65 to 2 grammes

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr., Ger, Jap, Russ, Swed and Swiss

Tests —Antipyrine Salicylate melts at 92°C ($197\ 6^{\circ}\text{F}$). The *P G* states 91° to 92°C ($195\ 8^{\circ}$ to $197\ 6^{\circ}\text{F}$), the *Fr Codex* 91°C ($195\ 8^{\circ}\text{F}$). A saturated aqueous solution affords, on the addition of a few drops of Fuming Nitric Acid, a green coloration, and with Tannic Acid Solution, a white precipitate, with Ferric Chloride T S a deep red coloration, changing, when largely diluted, to a reddish violet colour. 0.5 of a gramme mixed in 15 c c of Water and heated with 1 c c of Hydrochloric Acid affords a clear, colourless solution, from which fine white needles separate on cooling, which, when separated, washed with Water and dried, possess the m p and conform to the tests given under Acidum Salicylicum. A saturated aqueous solution of the salt shall not be affected by Hydrogen Sulphide, when warmed with Sodium Hydroxide Solution, and again warmed after the addition of Chloroform, no disagreeable odour of Phenyl isocyanide should be developed. It should leave no weighable residue when ignited with free access of air.

TOLYPYRIN —A body allied to Antipyrine (Phenazone), readily soluble in Water, and in Alcohol (90 p c), insoluble in Ether.

Antipyretic and analgesic, has been given in acute rheumatism —*L* '94, ii 991, *Pr* 1 383. See also under 'Feripyryn'.

Dose —5 to 20 grains = 0.32 to 1.3 grammes.

Tolysal (Tolypyrin Salicylate), sparingly soluble in Water, has been given in similar doses.

Tussol (Antipyrine Amygdalate) —In white granular crystals. Dose, for whooping cough in young children, 1 to 2 grains, older children may take as much as 7 grains. It should not be taken with Milk —*L* '95, i 1452, *P J* (3) xxv 912, 958.

PHENOL

See ACIDUM CARBOLICUM

Not Official

PHENOLPHTHALEIN

DI HYDROXY DIPHENYL METHYLIDE DIHYDROXY PHTHALOPHENONE

$\text{C}_{20}\text{H}_{14}\text{O}_4$, eq 315.72

In small, odourless crystals, or as a pale yellowish white powder, almost insoluble in Water, readily soluble in Alcohol (90 p c). For a long time it has been used as an indicator of neutrality in volumetric analysis, and is of interest from the magnificent pink coloration which its solution acquires by the action of alkalis. The weakest acids, on the other hand, destroy the colour. In recent years it has been found to possess a distinct aperient action, and has been introduced as an aperient under various names, Puigen, Aperione, etc. It may be administered in tablet form in doses of from 1 to 5 grains, and as a lozenge with chocolate basis containing 2 grains. If given to patients who suffer from hæmorrhoids a few doses have been shown (*B M J* '05, i 302) to produce an attack, and if the piles are troublesome at the time, they are aggravated, and often bleed freely.

Tests —Phenolphthalein melts at from 250° to 253°C (482° to $487\ 4^{\circ}\text{F}$). It dissolves readily in Alcohol with the formation of a colourless solution, which is neutral in reaction. The addition of Potassium or Sodium Hydroxide Solution to an alcoholic solution affords a magnificent pink coloration, which is destroyed on the addition of a slight excess of acid. The pink coloration produced with Sodium or Potassium Hydroxide Solution is permanent until the alkali Hydroxide

is converted into a neutral salt and the neutralising acid preponderates, it is therefore employed largely as an indicator of neutrality, and is the most trustworthy indicator for the determination of organic acids. The pink coloration produced by an alkali Hydroxide is destroyed by boiling with powdered Zinc, it is also destroyed by moist Carbonic Anhydride. Ammonia affords at first a pink coloration, but the colour is by no means permanent, and the substance does not therefore accurately indicate the point of neutralisation of Ammonia with an acid. 0.5 of a gramme when ignited with free access of air should leave no weighable residue.

NOSOPHEN (Tetraiodophenolphthalein)—A yellow, odourless powder, insoluble in Water, soluble in Ether and in Chloroform. Introduced as a substitute for Iodoform.

Dose—5 grains = 0.32 gramme

ANTINOSIN (Sodium Tetraiodophenolphthalein)—Blue prismatic crystals or blue amorphous powder, soluble in Water and in Alcohol (90 p.c.). Antiseptic. A substitute for Iodoform.

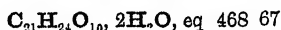
Dose—5 grains = 0.32 gramme

EUDOXIN (Bismuth Tetraiodophenolphthalein)—A reddish-brown, odourless powder, insoluble in Water. Introduced as a gastric and intestinal antiseptic.

Dose.—3 to 8 grains = 0.2 to 0.52 gramme.

Not Official

PHLORIDZIN



A glucoside, obtained from various Rosaceous trees.

A light crystalline powder, whitish, or pale yellow, slightly soluble in Water, 1 in 4 of Alcohol (90 p.c.).

A crystalline principle obtained from the bark of the stem and the root of the Cherry and some other allied trees.

It possesses the property of inducing artificial glycosuria in man and animals to which it is administered (*B.M.J.* '04, ii 890), and is of value as a test of renal inadequacy, but it is, of course, necessary to determine whether Glucose is already present in the urine, as if it is the test is valueless. The test consists in injecting subcutaneously 5 mg. of Phloridzin. The aid of Sodium Carbonate, in 20 to 80 minims of Water, immediately after the patient has emptied the bladder. If the kidneys are adequate, Glucose should appear in the urine in half an hour, when the amount may be estimated by Fehling's or by Pavv's method.

Dose—5 to 15 grains = 0.32 to 1 gramme, in mixtures, or in pills with 'Diluted Glucose'.

Official in Mex. (*Floricidina*)

Tests—Phloridzin loses at 100° C (212° F) its Water of crystallisation, which is theoretically equivalent to 7.6 p.c. According to Von Hager it melts at 107° C (224.6° F), again solidifies at 130° C (266° F), and again melts at 170° C (338° F) assuming at 200° C (398° F) a red colour, being decomposed with the formation of Prufin. When heated with dilute mineral acids it is decomposed, and the neutralised solution affords with Potassium-cupric Tartrate Solution a red precipitate. It dissolves in concentrated Sulphuric Acid with the production of a yellow colour, which turns red at a temperature between 27° and 50° C (77° and 122° F). When dissolved in an excess of Ammonia Solution and kept in contact with the air it gradually develops a violet or blue coloration. When heated with free access of air it should leave no weighable residue.

PHOSPHORUS.

PHOSPHORUS

P, eq 30 80

Fr, PHOSPHORE BLANC, Ger, PHOSPHOR, Ital, FOSFORO,
Span, FOSFORO

A colourless, or pale yellowish, almost translucent, waxy solid, having a characteristic, disagreeable odour. It rapidly oxidises on exposure to the air, and should be preserved under the surface of Water in well-stoppered bottles, away from the light and in a cool place. In the air it is luminous in the dark.

Solubility—Slightly soluble in Absolute Alcohol, 1 in 200 of Ether, 1 in 25 of Chloroform, 2 in 1 of Carbon Bisulphide, about 1 in 60 of Olive Oil, 1 in 60 of Oil of Turpentine, also in melted fats. Insoluble in Water.

Medicinal Properties—Given, but with doubtful success, as a nerve tonic, as an aphrodisiac, in rickets and in osteomalacia. Its prolonged use affects the structure of bones, causing them to become more dense, it also affects the liver and kidneys, leading to fatty degeneration. The preparations are *Oleum* and *Pilula*, and it has been combined with Cod-Liver Oil and other menstua, should be given with caution, as gastritis may be set up.

Sodium and Calcium Hypophosphites are forms of giving loosely-combined Phosphorus.

Dose, in pill or solution— $\frac{1}{100}$ to $\frac{1}{20}$ grain = 0.0006 to 0.0013 gramme.

Ph Ger maximum single dose, 0.001 gramme, maximum daily dose, 0.003 gramme.

Prescribing Notes—Generally given in pill form, to which may be added other tonics, such as Iron, Quinine and Strychnine, also dissolved in Almond Oil and Cod Liver Oil.

It should always be handled with caution, and be cut under Water.

Official Preparations—*Oleum Phosphoratum* and *Pilula Phosphori*. Used in the preparation of *Acidum Phosphoricum Concentratum* and *Calcii Hypophosphis*.

Not Official—*Elixir Phosphori*, *Elixir Phosphori Compositum*, *Pilula Phosphori cum Sevo*, *Pilula Phosphori Compositae*, *Pilula Phosphori cum Quinina*, *Sevum Phosphoratum*, *Spiritus Phosphori*, and *Tinctura Phosphori Composita*.

Antidotes—Stomach tube, emetics. Copper Sulphate is both emetic and antidote. 8 grains dissolved in Water every 5 minutes till vomiting is induced, then continue it in 1 grain doses every quarter of an hour, with 10 drops of Solution of Morphine if rejected, also 30 drops of old or French Oil of Turpentine every half-hour. Half an oz of Epsom Salts as a purgative. Demulcent drinks, but avoid oils and fats.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Ital, Jap, Mex (Fosforo), Norw, Port, Russ, Span, Swed, Swiss and U.S. Not in Hung.

Tests.—Phosphorus melts under Water at 44° to 45° C (111.2° to 113° F), the *B.P.* states that it melts at 43.3° C. (110° F). The

USP., *Fr Codex* and *PG* state 44° C (111 2° F) Its sp gr is given in the *BP* as 1 77, presumably at 15 5° C (60° F), the *USP* as 1 830 at 10° C (50° F) and 1 820 at 25° C (77° F.), and the *Fr Codex* as 1 83 When warmed in the air to a temperature a little over its mp, it takes fire and burns, producing dense white fumes which, when dissolved in Water, afford a solution possessing an acid reaction, which yields with Ammonium Molybdate Solution and Nitric Acid a lemon-yellow precipitate soluble in Ammonia Solution, and reprecipitated as a white crystalline precipitate on the addition of Magnesium Ammonio-sulphate Solution When oxidised with Nitric Acid it produces a solution yielding a similar precipitate to the above with similar reagents When dissolved in Carbon Bisulphide and poured on to a strip of filter paper, the latter, on the evaporation of the solvent, is instantly ignited It is required by the *USP* to contain not less than 99 5 pc of pure Phosphorus, but no method by which this percentage may be ensured is given

The two chief impurities are Arsenic and Sulphur, which are tested for after the oxidation of the Phosphorus by means of Nitric Acid The *BP* effects the oxidation by boiling 1 or 2 grammes of Phosphorus with 5 or 10 cc of Nitric Acid, mixed with an equal volume of Water The *USP* employs 1 gramme of Phosphorus, and digests it at a gentle heat on a water-bath with a mixture of 10 cc of Nitric Acid and 10 cc of Water, a current of Carbonic Acid gas being passed over the surface of the liquid whilst solution is being effected The *BP* requires that the resulting solution should yield no characteristic reaction with the tests for Arsenic, and only the slightest reactions with the tests for Sulphates The *USP* requirements are a good deal more definite, the solution, after the oxidation of the Phosphorus, is evaporated until no further nitrous vapours are evolved, and diluted with Water to 100 cc, 1 cc of which solution should not respond to the modified Gutzzeit's test for Arsenic, the presence of the latter much in excess of 1 in 100,000, is manifested by the formation of a distinct yellow-orange spot The addition of Barium Chloride TS to the remainder of the liquid is required to afford not more than a slight opalescence Phosphorus should dissolve readily and completely in Carbon Bisulphide Solution to form a clear solution, but the greatest caution is necessary in handling such solution, as the dissipation of the solvent is followed by immediate ignition

Preparations.

OLEUM PHOSPHORATUM.—PHOSPHORATED OIL.

1 of dry Phosphorus dissolved in 99 (by weight) of Almond Oil at 180° F (82 2° C) The Oil must first have been heated to 300° F. (149° C) for 15 minutes, cooled and filtered (about 1 in 100)

Dose.—1 to 5 minims = 0·06 to 0 3 c.c.

Foreign Pharmacopœias.—Official in Austr, 1 in 1000 Almond Oil, Belg and Span, 1 in 100 Almond Oil, Ital, 1 in 100 Olive Oil, Fr and Swed, 1 in 100 Almond Oil and Ether, Mex (*Acute fosforado*), 1 in 100 Sesame Oil,

Swiss, Phosphorus 1, Almond Oil 96, Alcohol (90 p c) 3, Natrium Sulphuricum Siccum 5 Not in the others

PILULA PHOSPHORI —PHOSPHORUS PILL

Dissolve 10 grains of Phosphorus in about 33 minims of Carbon Bisulphide, and add it to a mixture of 125 grains of melted White Beeswax and 125 grains of Lard cooled to a cream-like consistence, mix thoroughly, adding also 115 grains of Kaolin

The pill mass should be kept under Water in a light-proof stoppered bottle and, when required, it should be made into pills with one-third of its weight of powdered Gum Acacia and varnished

The finished pill is now 1 in 50, which is twice the strength of *B P* 1885

Dose —1 to 2 grains = 0.06 to 0.13 gramme

Foreign Pharmacopœias —Official in U.S., about $\frac{1}{100}$ grain of Phosphorus in each pill Not in the others

Not Official

ELIXIR PHOSPHORI —Compound Tincture of Phosphorus 1, Glycerin 4, should be preserved from the light Each fl. drin contains $\frac{1}{20}$ grain = 0.0018 gramme of Phosphorus —*B P C Formulary* 1901

Dose —15 to 60 minims = 0.9 to 3.6 c c

This has been incorporated in the *B P C* with the synonym **Syrupus Phosphori**, Syrup of Phosphorus

It should be freshly prepared as required

ELIXIR PHOSPHORI —Spirit of Phosphorus (*N F*), 21, Oil of Anise, 0.2, Glycerin, 56, Aromatic Elixir (*U S P*), *q s* to make 100 —*U S N F*

Elixir Phosphori Compositum *Syn* Syrupus Phosphori Composita —Compound Tincture of Phosphorus, 20 Oil of Anise, 0.20, Glycerin, 50, Aromatic Elixir, *q s* to produce 100 —*B P C*

Spiritus Phosphori *Syn* Tincture of Phosphorus —Phosphorus, 1.2, Absolute Alcohol (*U S P*), *q s* to make 1000 —*U S N F*

PILULA PHOSPHORI CUM SEVO —(1) Phosphorus, 10 grains, Mutton Suet, 90 grains, Purified Carbon Bisulphide, 40 minims Dissolve the Phosphorus in the Carbon Bisulphide, and incorporate with the Suet, previously rubbed into a smooth paste (2) Starch, 60 grains, Powdered Liquorice Root, 60 grains, Powdered Soap, 40 grains, Powdered Tragacanth, 12 grains, Glycerin, 48 minims Make into a pill mass

No 1 should be kept in a stoppered bottle, and incorporated with No 2 as required for dispensing 1 part of No 1 with 8 parts of No 2

They should be freshly prepared as required

Each 8 grain pill will contain $\frac{1}{8}$ grain of Phosphorus

PILULÆ PHOSPHORI COMPOSITÆ —Phosphorated Suet, 5, Quinine Sulphate, 12.50, Reduced Iron, 75, Strychnine, 0.50, Chloroform, 10, Compound Powder of Tragacanth, 5, Mucilage of Acacia, *q s* in 100 parts —*B P C*

This formula closely resembles that previously published in *Martindale*, except that 2 of these pills represent 1 of *Martindale's*

PILULÆ PHOSPHORI CUM QUININA —Phosphorated Suet, 10, Quinine Sulphate, 50, Chloroform, 20, Compound Powder of Tragacanth, 10, Mucilage of Acacia, *q s* in 100 parts —*B P C*

This formula closely resembles that previously published in *Martindale*

SEVUM PHOSPHORATUM (10 p c) —Phosphorus, 1, Pure Carbon Bisulphide, 5, Dissolve, and add Prepared Suet, 9 Add a little of the Suet at first, mix quickly, add the remainder, mix thoroughly, and allow the Bisulphide to evaporate —*Martindale*

This has been incorporated in the *B P C*

TINCTURA PHOSPHORI COMPOSITA —Dissolve 12 grains Phosphorus in $2\frac{1}{2}$ fl. oz Chloroform by the aid of a gentle heat, add the solution to

12½ fl oz Ethylic Alcohol and shake well. Should be preserved in well stoppered bottles and kept from the light.

10 minims contain $\frac{1}{10}$ grain of Phosphorus

Dose—3 to 12 minims = 0.18 to 0.71 c.c.

BPC Formulary 1901, incorporated in the *BPC* with a slight increase in strength, as follows—

Phosphorus, 0.20, Chloroform, 17, Absolute Alcohol, q.s. to produce 100

PHYSOSTIGMATIS SEMINA.

CALABAR BEAN

FR, FÈVE DU CALABAR, GER, KALABARBOHNE, ITAL, FAVA DEL CALABAR, SPAN, HABA DEL CALABAR

The ripe Seeds of *Physostigma venenosum*, Balfour

The Seeds official in the *BP* are not required to yield any definite amount of alkaloids. The *USP* Seeds are required to yield not less than 0.15 p.c. of Ether-soluble alkaloids. The Seeds are not official in the *PG*.

— Indigenous to Western Africa

The chief constituent is a poisonous crystalline alkaloid, **Physostigmine** or **Eserine**

Medicinal Properties.—Myotic, antispasmodic. It increases the flow of saliva and most of the other secretions. Used in tetanus, but its principal use is in ophthalmic work. See '**Physostigminæ Sulphas**'.

Official Preparation.—*Extractum Physostigmatis*. Used to prepare *Physostigmatis Sulphas*.

Not Official—*Tinctura Physostigmatis*

Foreign Pharmacopœias—Official in Belg (*Semen Calabariense*), Jap (*Semen Physostigmatis*); Mex (*Haba de Calabar*), Port (*Favo do Calabar*), Span (*Haba del Calabar*), Swed (*Semina Calabar*), US (*Physostigma*). Not in the others.

Descriptive Notes—Calabar Beans as recently imported are rather smaller and browner in colour than formerly. Formerly the seeds were almost black in colour and about $1\frac{3}{8}$ in (34 mm) long, $\frac{3}{4}$ in (19 mm) broad, and $\frac{1}{2}$ to $\frac{5}{8}$ in (12.5 to 15 mm) in thickness, but the dimensions given in the official description are 'usually about 1 in long (25 mm), $\frac{3}{4}$ in (18 mm) broad, and $\frac{1}{2}$ in (12 mm) thick'. The hilum extends nearly the whole length of the curved margin of the seed, which is elongate reniform in outline. The seed coat is hard, dark reddish-brown, and slightly rough. The two seeds connected have a cavity between them. The seed has neither taste nor odour. At one time a seed of a different species, nearly cylindrical and scarcely curved, but about the same size, was offered in London as Calabar Bean. It has been referred to *Physostigma cylindrospermum*, Holmes. Other seeds quite different in shape and size from Calabar Bean have been offered as substitutes in the drug market, but none of these could be mistaken for the genuine drug.

Tests.—Calabar Bean is assayed in the *U S P* by the following process —A weighed quantity of 20 grammes is introduced into an Erlenmeyer flask of about 250 cc capacity and shaken well during 10 minutes with 200 cc of Ether, 10 cc of a 1 in 20 aqueous Sodium Bicarbonate Solution is added and the mixture shaken vigorously during 4 hours. A measured quantity of 100 cc (= 10 grammes of Calabar Bean) of the Ether solution is decanted into a separator after the powder has been allowed to settle. Sufficient Normal Volumetric Sulphuric Acid Solution and is added to render the liquid acid, and 10 cc of Water. The extraction is repeated first with a mixture of 2 cc of Normal Volumetric Sulphuric Acid Solution and 8 cc of Distilled Water, and then with a mixture of 1 cc of Normal Volumetric Sulphuric Acid Solution and 9 cc of Water. The acid liquids are in each case separated and transferred to another separator. Sufficient of a 1 in 20 aqueous Sodium Bicarbonate Solution is added to the combined acid liquids in the separator to render them alkaline to red Litmus paper, and the liberated alkaloids are extracted by shaking with 3 successive quantities each of 25 cc, 20 cc, and 15 cc of Ether, the ethereal solutions being separated in each instance, mixed, transferred to a flask, and carefully evaporated on a water bath. The residue when dry is dissolved in 5 cc of Tenth-normal Volumetric Sulphuric Acid Solution, 20 cc of absolutely neutral Ether added, and the mixture transferred to a stoppered bottle, rinsing the flask with 80 cc of Water. The excess of Tenth-normal Volumetric Acid Solution is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, 5 drops of Iodeosin T S being used as an indicator of neutrality. The number of cc of Fiftieth-normal volumetric alkali solution required divided by 5, the quotient subtracted from 5, the difference multiplied first by 0.0273 and then by 10, gives the percentage of Ether-soluble alkaloids contained in the Calabar Beans.

Preparation

EXTRACTUM PHYSOSTIGMATIS —**EXTRACT OF CALABAR BEAN**
16 of Calabar Bean treated with 80 of Alcohol (90 p c), the liquid evaporated to a soft extract and mixed with 3 times its weight of Milk Sugar to form a firm extract.

It is about one fourth the strength of *B P* 1885, and of the Foreign Pharmacopœias.

Dose — $\frac{1}{4}$ to 1 grain = 0.016 to 0.032 gramme

As it does not form a clear solution with Water, such solution should be filtered.

Foreign Pharmacopœias —Official in Jap, Port and U S. Not in the others.

28 lb of Calabar Beans, treated with Alcohol (90 p c), yielded 2.07 p c of extract, this extract yielded 5.74 p c of alkaloids, which is equal to nearly 0.12 p c of alkaloids in the Beans.

The same powder treated with boiling Alcohol (90 p c) in an exhaustion apparatus yielded 4.66 p c of extract, which extract yielded 3.2 p c of alkaloids, which is equal to nearly 0.15 p c of alkaloids in the Beans.

The extract of Calabar Bean official in the *B P* is not a

standardised preparation. The *USP* extract is required to contain 2 p c of Ether-soluble alkaloids, the extract is not official in the *PG*.

Tests.—The *USP* employs a process for the assay of the extract of which the following contains the essential features.—A weighed quantity of 1 gramme of the extract is transferred to a small porcelain evaporating basin and digested for 5 minutes at a temperature below the boiling point of boiling Water with 5 c c of Alcohol (48.9 p c). After the addition of 5 grammes of very clean fine sand the mixture is evaporated to dryness on a water-bath and triturated thoroughly to ensure a uniform mixture, the contents of the dish are transferred as soon as dry to an Erlenmeyer flask (using a little more clean sand to transfer the residue to the flask) and shaken with 100 c c of Ether, a measured quantity of 10 c c of a 1 in 20 aqueous Sodium Bicarbonate Solution is added and the contents vigorously shaken at intervals for 1 hour. When the powder has settled, 50 c c of the Ether solution is decanted into a separator, and sufficient Normal Volumetric Sulphuric Acid Solution to render the liquid acid in reaction towards blue Litmus paper, and 10 c c of Distilled Water are added and the liquid shaken, the shaking being repeated with 2 successive quantities each of 2 c c of Normal Volumetric Sulphuric Acid solution and 8 c c of Distilled Water, and 1 c c of Normal Volumetric Sulphuric Acid Solution and 9 c c of Distilled Water, the acid liquids being in each case separated and transferred to a second separator. Sufficient of a 1 in 20 aqueous Sodium Bicarbonate Solution is added to the mixed acid liquids to render them alkaline in reaction towards red Litmus paper, and the liberated alkaloids are extracted by agitation with 3 successive quantities each of 25 c c, 20 c c, and 15 c c of Ether, the ethereal solution being in each instance separated and transferred to a flask. They are mixed, evaporated on a water-bath, and when dry the residue is dissolved in 2 c c of Tenth-normal Volumetric Sulphuric Acid Solution, and when dissolved is transferred to a 200 c c flask, washing the flask with Water and adding enough Water to bring the volume to about 90 c c. 25 c c of Ether is added and the excess of Tenth-normal Volumetric Acid Solution is titrated with Fiftieth-normal Volumetric Potassium Hydroxide Solution, using 5 drops of Iodeosin TS as an indicator of neutrality. The number of c c of Fiftieth-normal Volumetric Potassium Hydroxide Solution required divided by 5, the quotient subtracted from 2, and the difference multiplied first by 0.0273 and then by 200, yields the percentage of Ether-soluble alkaloids present in the sample operated on.

Not Official

TINCTURA PHYSOSTIGMATIS—Calabar Bean, 1, Alcohol (90 p c), 5.

Dose—5 to 15 minims = 0.3 to 0.9 c c

BPC Formulary 1901, is incorporated in the *BPC*.

Foreign Pharmacopœias—Official in *US*, 1 in 10. Not in the others. The *USP* tincture is required to contain 0.014 p c w/v of Ether-soluble alkaloids. Neither the *BP* nor the *PG* includes a tincture of Physostigmine.

Tests—A measured quantity of 100 c c of the tincture is evaporated to dryness on a water bath, and the proportion of Ether soluble alkaloid from Physostigmine in the extract so obtained is determined by the *USP* process given under *Extractum Physostigmatis*, and when calculating the result of the volumetric determination into terms of Ether soluble alkaloids the product should be multiplied by 2 instead of 200, that is to say, the number of c c of Fiftieth-normal Volumetric Potassium Hydroxide Solution required to neutralise the excess of Tenth normal Volumetric Acid Solution divided by 5, the quotient subtracted from 2, the difference multiplied first by 0.0273 and then by 2, yields the percentage w/v of Ether soluble alkaloids in the sample operated on

PHYSOSTIGMINÆ SULPHAS.

PHYSOSTIGMINE SULPHATE

B P Syn —ESERINL SULPHAT

$(C_{15}H_{21}N_3O_4)_2, H_2SO_4$, eq 643.80

FR, SULFATE D'ESERINE, GER, PHYSOSTIGMINSULFAT, ITAL, ESERINA SOLFATO, SPAN, SULFATO DL ESERINA

A whitish or yellowish-white, very deliquescent micro-crystalline powder, possessing a bitter taste. It should be carefully preserved in small, well-stoppered glass bottles of a dark amber tint, or in sealed tubes protected from the light.

It is the Sulphate of an alkaloid obtained from Calabar Bean. The Sulphate alone is official in the *B P*. The *P G* and the *USP* include both the Sulphate and the Salicylate. The *Ph Codex* only includes the Salicylate. The Pharmacopœia formula for the Sulphate is given as xH_2O . The *USP* gives the formula for the anhydrous salt. *P G* does not include formulas.

Solubility.—4 in 1 of Water, $2\frac{1}{2}$ in 1 of Alcohol (90 p c)

Medicinal Properties—It is used to contract the pupil in ciliary paralysis due, *eg*, to diphtheria, to reduce intra-ocular tension in glaucoma, etc, to prevent or reduce prolapse of the iris after corneal wounds, to diminish the amount of light in painful affections of the eye, to break down adhesions due to iritis, its use being alternated with that of Atropine, and to remove the prolonged dilatation and paralysis produced by the latter. In tetanus it is to be given fearlessly, $\frac{3}{10}$ grain hypodermically frequently repeated, the patient being carefully watched. An antidote in Strychnine poisoning.

Dose— $\frac{1}{10}$ to $\frac{1}{10}$ grain = 0.0011 to 0.0032 gramme

Dutch maximum single dose, 0.001 gramme, maximum daily dose, 0.003 gramme

Prescribing Notes—The salts of Physostigmine as well as the solutions are liable to become pink by oxidation. They should be kept in yellow non-actinic glass bottles, and as much as possible preserved from the air.

If desirable the coloration of the solutions may be prevented by the addition of a trace of Hypophosphorous Acid or Sulphurous Acid.

Official Preparation—Lamellæ Physostigminae

Not Official.—Guttæ Physostigminae, Guttæ Physostigminae cum Cocaina, Physostigmina, Unguentum Physostigminae, Physostigminae Hydrobromidum, and Physostigminae Salicylas

Foreign Pharmacopœias.—Official in Dutch, Ger., Jap., Mex. (Sulfato de Eserina), and in others.

Tests.—Physostigmine Sulphate is stated by the *U.S.P.* to soften at 130°C (266°F), and to melt at 140°C (284°F), neither the *B.P.* nor the *P.G.* makes any reference to the m.p. It dissolves readily in Water, yielding a solution which is neutral in reaction towards Litmus paper. The *U.S.P.* states that it possesses an acid reaction towards blue Litmus paper. The addition of Potassium or Sodium Hydroxide Solution causes a white precipitate of the alkaloid which quickly turns pink, and the precipitate itself dissolves in an excess of the Hydroxide Solution, with the production of a pinkish or red coloured solution, when evaporated to dryness with Ammonia Solution a bluish coloured residue remains, and if this residue be dissolved in very dilute acid the solution assumes a red coloration by reflected light, and a blue coloration by transmitted light. A trace of the salt dissolved in Fuming Nitric Acid yields a yellow solution, which, when evaporated to dryness on a water-bath, yields a residue of a green colour. The *U.S.P.* states that 5 mg. of the salt yield a yellow coloured solution when dissolved in Nitric Acid, on heating this solution the colour changes from orange to blood-red, and leaves on evaporation to dryness a green-coloured residue. The residue on exposure to the fumes of Nitric Acid yields a violet-blue colour, and on the addition of a drop of Nitric Acid a reddish-violet coloured solution changing rapidly to blood-red, and on standing, or on dilution, a greenish-yellow. The most distinctive test for Eserine is the marked mydriasis which it produces on the pupil of the eye, a highly diluted aqueous solution applied to the conjunctiva causes almost immediate contraction of the pupil. Sulphuric Acid yields only a faint yellow colour with the salt. Sulphuric Acid containing a crystal of Potassium Iodate yields a pale purple coloration rapidly changing to yellowish-red. The aqueous solution, with or without acidification with Hydrochloric Acid, yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. The Sulphate may be distinguished from the Salicylate by yielding when dissolved in Water a yellowish-white precipitate with Platonic Chloride Solution, and by the non-production of a violet colour on the addition of Ferric Chloride T.S. When ignited with free access of air it should leave no weighable residue.

Preparation.

LAMELLÆ PHYSOSTIGMINÆ. DISCS OF PHYSOSTIGMINE
Gelatin discs, containing $\frac{1}{100}$ grain of Physostigmine Sulphate

Now made with Physostigmine Sulphate instead of Physostigmine

Foreign Pharmacopœias.—Official in Ital., Dischi Oftalmici con Eserina, etc. (0.001 gramme Eserine Salicylate). Not in the others.
Hypodermic lamellæ containing $\frac{1}{100}$ grain.—*Bartholomew's*

Not Official

GUTTÆ PHYSOSTIGMINÆ.—Physostigmine Sulphate, $\frac{1}{2}$, 1, 2, or 4 grains, Water, 1 fl. oz.—*London Ophthalmic*

GUTTÆ PHYSOSTIGMINÆ CUM COCAINA—Physostigmine

Sulphate, 1 grain, Cocaine Hydrochloride, 4 grains, Water, 1 fl oz—*London Ophthalmic*

PHYSOSTIGMINA Eserine $C_{15}H_{21}N_3O$, eq 273.23—Colourless or pale pink glistening crystals, very slightly soluble in Water, readily soluble in Alcohol (90 p c), Ether and Chloroform. It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the air and light.

It was official in *BP* '85, but deleted from *BP* '98, the Sulphate being made official.

Unguentum Physostigminæ (Unguentum Eserinæ) Physostigmine, 1 grain, Soft Paraffin, 1 oz, is given in *London Ophthalmic*.

Tests—Physostigmine melts at 102° to 103° C (215.6° to 217.4° F). Petit and Polonovski give 105° to 106° C (221° to 222.8° F). It dissolves very slightly in Water, it is readily soluble in Alcohol and Ether, the solutions in these solvents being strongly levogyrate. The aqueous solution is alkaline in reaction towards red Litmus paper, and precipitates Ferric Hydroxide from Ferric Chloride Solution, provided the latter is not too acid, it answers the tests characteristic of Eserine given under Physostigmine Sulphate.

PHYSOSTIGMINÆ HYDROBROMIDUM—In colourless crystals, very soluble in Water.

PHYSOSTIGMINÆ SALICYLAS *Syn* Eserina Salicylas $C_{15}H_{21}N_3O \cdot C_7H_5O_2$, eq 410.24—Colourless or faintly yellowish acicular crystals, becoming coloured on exposure to light and air. Soluble 1 in 180 of Water, 1 in 15 of Alcohol (90 p c). The Salicylate of the *Fr Codex* contains 66.59 p c of Eserine and 33.41 p c of Salicylic Acid.

It is the Salicylate of Physostigmine, an alkaloid obtained from Calabar Bean. It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the light and air, as it rapidly acquires a reddish tint on exposure to their combined influence.

A sterilised solution containing 0.2 grammes of Eserine Salicylate in 40 grammes of Olive Oil is stated (*P J* '05, 1 589) to keep indefinitely and to produce immediate and painless action upon the eye.

The theory that the formation of Rubeserine is due to the absorption of atmospheric Ammonia by the solution is stated (*CD* '05, 11 515) to be inadmissible. Conjunctivitis has occasionally been traced to the use of oxidised solutions of Eserine. A reference is again made to the use of oily solutions of the alkaloid or of the Salicylate.

Ph Ger maximum single dose, 0.001 gramme, maximum daily dose, 0.008 gramme.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Fr., Ger., Hung., Ital., Jap., Mex., Norw., Russ., Swed., Swiss and U.S. Not in the others.

Tests—Physostigmine Salicylate melts at about 179° C (354.2° F). The *USP* states that it softens and assumes a slightly yellow colour at 160° C. (320° F) and melts at 178.9° C (354° F). *Fr Codex* gives 182° C (359.6° F). It dissolves in Water, forming a clear solution, which is faintly acid in reaction towards blue Litmus paper. It should answer the tests characteristic of Physostigmine given under Physostigmina Sulphate. Sulphuric Acid, containing a drop of Formaldehyde Solution in each c.c., produces a bright pink colour. Sulphuric Acid, with a few crystals of Cane Sugar, produces a yellow colour changing to brownish-purple and ultimately to greenish-black. An aqueous solution of the salt affords with Ferric Chloride T.S. a deep violet coloration. The addition of Bromine Water, if present in excess, produces an intense red coloration in the salt or its solution. It may be distinguished from the Sulphate by not yielding a precipitate on the addition of Platonic Chloride Solution, and by the production of a violet coloration with Ferric Chloride T.S. It should leave no weighable residue when ignited with free access of air.

Not Official
PHYTOLACCA.

Both the **Fruit** (Poke fruit) and the **Root** (Poke root) of *Phytolacca decandra*, L., are official in U S. The Root is official in Jap.

The Fluid Extract has been recommended for inflamed and painful mammae, internally and as a local application—*B M J* '87, 11 844. It has also been used in orchitis—*T G* '85, 622. In large doses it is emetic, purgative, and slightly narcotic.

FLUIDEXTRACTUM PHYTOLACCÆ RADICIS (U S)—1 fl oz is equal to 1 oz of root.

Dose.—As an alterative, 1 to 5 minims = 0.06 to 0.08 c c.

Official in Jap.

TINCTURA PHYTOLACCÆ—Poke Root, 1, Alcohol (45 p c), 10. Dose—3 to 10 minims = 0.18 to 0.6 c c—*Martindale*.

This has been incorporated in the *B P C*.

PHYTOLACCIN—An eclectic remedy used in rheumatic and syphilitic conditions. In pill as a cholagogue and alterative, $\frac{1}{4}$ to $\frac{1}{2}$ grain = 0.016 to 0.032 gramme, purgative, 2 to 4 grains = 0.13 to 0.26 gramme.

Not Official
PICRORHIZA

The dried Rhizome of *Picrorhiza Kurroa*, Royle, dose in powder, as a tonic, 10 to 20 grains = 0.65 to 1.3 grammes, as an antiperiodic, 40 to 50 grains = 2.6 to 3.2 grammes, is official in the *Ind* and *Col Add* for India and the Eastern Colonies, also **Extractum Picrorhizæ Liquidum**, 1 in 1 Fluid Extract made with Alcohol (60 p c), dose, 20 to 60 minims = 1.2 to 3.6 c c, and **Tinctura Picrorhizæ**, Picrorhiza 1, Alcohol (45 p c) 8, by maceration, dose, $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c c.

PICROTOXINUM.

PICROTOXIN

Colourless, shining, prismatic crystals, or micro-crystalline powder permanent in the air, possessing an intensely bitter taste. It is the neutral principle obtained from 'Cocculus Indicus' described in *B P* as the Fruits of *Anamita paniculata*, Colebr.

Picrotoxin is stated to be not a simple body but a compound, containing 34 p c of Picrotin and 66 p c of Picrotoxinin, but its composition cannot be considered as definitely settled.

Solubility—1 in 334 of Water, 1 in 13 $\frac{1}{2}$ of Alcohol (90 p c).

Medicinal Properties.—Anhydrotic, $\frac{1}{60}$ gram at bedtime has been given as a remedy against immoderate sweating in phthisis.

Externally used with caution as an ointment (8 grains to 1 oz) for pediculi.

Dose.— $\frac{1}{100}$ to $\frac{1}{25}$ gram = 0.0006 to 0.0024 gramme.

Antidotes—Stomach-tube, or emetic, Chloral, and Potassium Bromide.

Foreign Pharmacopœias—Official in Fr and Mex. Not in the others.

Tests.—Pure Picrotoxin melts at 199° to 200° C (390.2° to 392° F), the *B P* melting at 192.2° C (378° F), the *Fr Codex* at 200° C (392° F). It is officially stated to be soluble in 10 parts of

Potassium Hydroxide Solution, the liquid so formed producing a red precipitate with Potassio-cupric Tartrate (Fehling's solution). The test may also be applied to a cold saturated aqueous solution, 5 c c of which will give a distinct reaction, if a similar quantity of Pavy's solution be added and the liquid boiled the blue colour will completely disappear. It dissolves in Sulphuric Acid with the production of a bright yellow coloured solution, changing to orange-red on warming, and very gradually to reddish-brown. It dissolves in Nitric Acid, and the liquid on evaporation leaves a reddish-yellow residue, becoming bright red when moistened with Potassium Hydroxide Solution. It may be distinguished from alkaloids by not yielding when dissolved in Water precipitates with Platinic Chloride, Potassio-mercuric Iodide, Mercuric Chloride, Tannic Acid, Potassium Ferrocyanide and Ferricyanide Solutions, and most of the general reagents for alkaloids. When ignited with free access of air it should leave no weighable residue.

PILOCARPINÆ NITRAS.

PILOCARPINE NITRATE

$C_{11}H_{16}N_2O_2$, HNO_3 , eq 269 23

FR, AZOTATE DE PILOCARPINE, GTR, PILOCARPINITRAT,
ITAL, PILOCARPINA NITRATO

It is the Nitrate of Pilocarpine, an alkaloid obtained from Jaborandi Leaves. A synthetic Pilocarpine has also been produced.

Pilocarpine Nitrate occurs as white distinct crystals. Jowett states that the Nitrate is the most convenient to use in medicine, on account of its stability in the air, the Hydrochloride being hygroscopic in moist air. It should be kept in well-stoppered glass bottles of a dark amber tint and in a cool atmosphere.

Solubility—1 in 8 of Water, 1 in 50 of Alcohol (90 p c), almost insoluble in Ether and in Chloroform.

Medicinal Properties—A powerful diaphoretic and sialagogue. Is useful in the dropsy and thirst of Bright's disease, in uræmia, and to remove pleural and peritoneal effusion. It should be used with caution, if at all, in cardiac dropsy. It contracts the pupil, and has been used in detachment of the retina, glaucoma and intra-ocular hæmorrhage, it has been given in bronchitis and asthma, and in chronic poisoning by lead, arsenic or mercury. Useful in deafness due to disease of the auditory nerve. A good antidote in Belladonna poisoning.

It has been used to increase the growth of the hair as a **Lotion** containing 1 or 2 grains to 1 fl oz, and as an **Ointment** containing 4 to 8 grains to 1 fl oz.

In pneumonia, $\frac{1}{10}$ to $\frac{1}{8}$ grain hypodermically.—*L* '03, 1 1369, '03, 11 342, *B M J* '03, 11 788. $\frac{1}{4}$ to $\frac{1}{2}$ grain by the mouth.—*L* '03, 1 823.

In detachment of the retina, 1 cg of the Nitrate injected in a 2 p c solution, —*B M J E*, '99, 11 68.

$\frac{1}{2}$ grain injected subcutaneously in severe uræmia of Bright's disease — *Pr* lxvii 657

Objection taken to the *BP* dose, $\frac{1}{30}$ to $\frac{1}{2}$ grain, when given by the mouth as being too high. In one case $\frac{1}{30}$ grain caused vomiting every time it was taken, in another $\frac{1}{30}$ grain caused profuse sweating and exhaustion lasting some hours. Probably $\frac{1}{30}$ grain is the highest initial dose that should be given — *BMJ* '02, ii 1104

Its use in puerperal eclampsia has been abandoned at the Rotunda Hospital, Dublin — *L* '05, ii 749

Of great value in all forms of pruritus, but especially that of the vulva; $\frac{1}{8}$ to $\frac{1}{4}$ grain by mouth only when itching manifests itself, the addition of $\frac{1}{12}$ grain Atropine may sometimes be necessary to prevent sweating — *MR* '07, i 858.

While the most powerful of the internal diaphoretics, it must be used cautiously owing to its tendency to depress the heart — *BMJ* '06, ii 1450

Dose. — $\frac{1}{30}$ to $\frac{1}{2}$ grain = 0.0032 to 0.032 gramme

Prescribing Notes. — Most ^{sed by hypodermic injection, also} given in solution, and in pills with ^{and Glucose} Supplied also in hypodermic tablets, $\frac{1}{10}$, $\frac{1}{8}$, $\frac{1}{4}$ and $\frac{1}{2}$ grain

The nearly equal solubility of the Pilocarpine Nitrate and Pilocarpidine Nitrate allows them to crystallise together. With the Hydrochlorides the difference in solubility is much more marked, so that a Pilocarpine Hydrochloride can be obtained containing very little Pilocarpidine.

The Pilocarpine Hydrochloride is preferred in all other countries, see below, and in London is more frequently prescribed than the Nitrate, but it is incompatible with Silver salts, with which Pilocarpine is sometimes used.

Not Official. — *Guttæ Pilocarpinæ*, *Injectio Pilocarpinæ Nitratis*, *Pilocarpine*, *Pilocarpinæ Hydrochloridum*, *Pilocarpinæ Phenas*, and *Pilocarpinæ Salicylas*

Foreign Pharmacopœias. — Official in *Fr*, *Mex* and *US*. Not in the others. *Fr* and *Mex* have Pilocarpine.

Antidote. — Belladonna by the mouth, or Atropine hypodermically.

Tests. — Pilocarpine Nitrate is required by the *BP* to form with H_2SO_4 a yellowish solution, which on the addition of NaOH gradually assumes an emerald-green colour, the *USP* states that with Sulphuric Acid a colourless solution is produced. A characteristic reaction is the mydriasis produced by a dilute aqueous solution of the salt. The *BP* does not include a m.p., the *USP* states that it melts at 170.9°C (339.7°F). Pure Pilocarpine Nitrate, according to Jowett, melts at 173° to 178°C (343.4° to 352.4°F), according to Petit and Polonovski at 177° to 178°C (350.6° to 352.4°F), the *Fr Codex* (1908) gives 177°C (350.6°F). It has a specific rotation of $+80^\circ$ to $+88^\circ$. *Fr Codex* $+82.2'$ at 18°C . The addition of an excess of Ammonia Solution to an aqueous solution of the salt should not afford a precipitate, the addition of Sodium Hydroxide Solution to dilute aqueous solutions of the salt affords no precipitate, but if the solutions be sufficiently concentrated, a white turbidity is produced. The separated alkaloid should answer the tests distinctive of Pilocarpine under that heading. As an additional test Jowett has suggested the formation of a crystalline Picrate which should melt sharply at 147°C (296.6°F). When dissolved in Water it affords a clear colourless solution which should possess a faint acid reaction towards blue Litmus paper, and which,

when kept cool and mixed with an equal volume of Sulphuric Acid, affords a brown ring at the junction of the two liquids when a solution of Ferrous Sulphate is carefully poured on the surface of the mixture. Pilocarpine Nitrate may be distinguished from the Hydrochloride by rubbing the salt with an equal weight of Mercurous Chloride: in the case of the Nitrate no black coloration is produced, in the case of the Hydrochloride the Mercurous Chloride is reduced with the formation of metallic Mercury, a blackening in colour simultaneously occurring. According to the *USP* it may be distinguished from other alkaloids by dissolving 10 to 20 mg of the salt in 2 c.c. of Water, and adding 2 c.c. of a slightly acidified Hydrogen Peroxide Solution, and pouring upon the surface of the liquid a layer of Benzene. On the addition of 3 or 4 drops of a 1 in 300 Potassium Bichromate Solution the Benzene layer will acquire a violet colour; if the mixture be gently shaken, the aqueous layer remaining yellow. The *USP* states that if more than 20 mg be taken the Benzene turns blue, and the reaction is no longer characteristic. The *BP* and the *USP* require that when ignited with free access of air it shall leave no residue.

Not Official

GUTTÆ PILOCARPINÆ—Pilocarpine Nitrate, 2 grains, Distilled Water, 1 fl. oz.—*London Ophthalmic*

INJECTIO PILOCARPINÆ NITRATIS—Pilocarpine Nitrate, 1, Water, 20. Dose, 2 to 6 minims.—*London Ophthalmic*

Pilocarpine Nitrate, 1 grain, Distilled Water, 12 minims. Dose, 1 to 4 minims.—*Guy's*

To prepare the patient for the injection, remove the nightshirt, wrap him closely in a warm blanket, and cover him with two more blankets. Put hot-water bottles to his feet, and give him hot drinks freely. After the sweating has ceased, remove the blankets gradually, dry the skin thoroughly, and leave him between warm dry blankets.—*Guy's*

PILOCARPINE ($C_{11}H_{16}N_2O_2$, eq 206.65)—It is the principal alkaloid contained in Jaborandi Leaves, and may also be produced synthetically. It forms a colourless and odourless thick syrup, which becomes thinner on warming. It is readily soluble in Water, Alcohol (90 p.c.), and Chloroform. The solutions are dextrogyrate. It gives no colour reaction with strong Sulphuric or Nitric Acid, with Sulphuric Acid and Potassium Bichromate it gives a dark green coloration. When titrated with an excess of Calomel the latter is reduced to metallic Mercury, a darkening in colour resulting. It is precipitated by the usual alkaloidal reagents, such as Potassium mercuric Iodide (Mayer's) Solution, Iodo-potassium Iodide (Wagner's) Solution, Tannic Acid, etc. It is stated by Allen to give no reaction with Picric Acid, but Jowett (*YBP* '99, 436) states that the alkaloid affords a yellow precipitate which dissolves on warming, again separating out in needles when the solution cools. It may be determined by titration with Tenth-normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Iodæosin Solution as an indicator of neutrality. 1 c.c. of Tenth-normal Volumetric Acid Solution is equivalent to 0.02066 gramme of pure Pilocarpine.

PILOCARPINÆ HYDROCHLORIDUM Pilocarpine Hydrochloride $C_{11}H_{18}N_2O_2 \cdot HCl$, eq 242.84—Colourless or white cubical crystals, deliquescent in moist air, soluble in less than its own weight of Water, 1 in 10 by weight of Ethyl Alcohol, almost insoluble in Ether or Chloroform.

A more definite salt than the Nitrate, being more easily separated from accompanying Hydrochlorides of the other bases, but deliquescent in moist air.

It should be kept in well-stoppered glass bottles of a dark amber tint and in

a cool atmosphere, it should also be kept as far as possible from contact with a moist atmosphere

Dose — $\frac{1}{2}$ gr to $\frac{1}{2}$ grain = 0.032 to 0.033 gramme

Ph Ger maximum single dose, 0.02 gramme, maximum daily dose, 0.04 gramme

Incompatibles.—Alkalis, and Alkaline Carbonates, Lead, Mercurous and Silver salts

Foreign Pharmacopœias —Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw (*Chloietum Pilocarpicum*), Russ, Swed, Swiss and U S Not in the others

Tests —Pure π is that officially adopted when anhydrous, according to Petit and Polonovs π is that officially adopted in the *Fr Codex* (1908) The *U S P* gives the m.p. of the salt after drying for several hours at 100° C (212° F) as 195.9° C (384.5° F) *P G* gives the m.p. as 198° to 195° C (379.4° to 383° F) Jowett states (*Y B P* '99, 441) that when the salt dried at 100° C (212° F) is heated in a capillary tube it melts at 200° to 204° C (392° to 399.2° F) Solutions of the salt are dextrogyrate, the specific rotatory power of the aqueous solution being +80° to +92° *Fr Codex* gives +91° at 18° C for the 2 p.c w/v aqueous solution When dissolved in Water it forms a clear colourless solution which should have a neutral or at the most but faintly acid reaction towards Litmus paper It should dissolve in Sulphuric Acid to form an almost colourless liquid, Hydrochloric Acid gas being simultaneously evolved On the addition of a tiny crystal of Potassium Bichromate a bright emerald-green coloration is produced Ammonia Solution should not produce a precipitate when added to a concentrated aqueous solution of the salt, and Sodium or Potassium Hydroxide Solution added to a similar solution should produce but a few oily droplets The separated alkaloid should answer the tests given under that heading The aqueous solution should yield when acidified with Nitric Acid, and on the addition of Silver Nitrate Solution, a white curdy precipitate insoluble in Nitric Acid, soluble in Ammonia Solution or in Potassium Cyanide Solution It may be distinguished from the Nitrate by yielding a black coloration when rubbed with an equal quantity of Mercurous Chloride Pilocarpine Nitrate under similar conditions yields no black coloration It may be distinguished from other alkaloids by the Ammonia test given above or by dissolving 10 to 20 mg of the salt in 2 c.c of Water, mixing the solution with 2 c.c of a slightly acidified Hydrogen Peroxide Solution, carefully adding sufficient Benzene to form a small layer on the surface of the liquid, and adding 3 or 4 drops of a 1 in 300 Potassium Bichromate Solution, on gently shaking the mixture the Benzene layer will acquire a violet colour, the aqueous layer remaining yellow

The *U S P* states that if more than 20 mg be used the Benzene turns blue and the reaction is no longer characteristic

When ignited with free access of air it should leave no residue

It is official in 15 out of the 17 Foreign Pharmacopœias

PILOCARPINÆ PHENAS —A colourless, oily liquid, soluble in Water and in Alcohol, has been recommended in phthisis and intermittent fevers, 1 fl dr of a solution of 1 grain in 10 fl oz of 2 p.c Carbolic Acid Solution (*Aseptoline*) injected into the abdominal wall —*P J* '96, ii 379, '98, i 84

PILOCARPINÆ SALICYLAS.—Colourless crystals, or as a white crystalline powder, soluble in Water, less soluble in Alcohol (90 p.c) Employed for purposes similar to those of the Nitrate or Hydrochloride

It should be kept in well-stoppered glass bottles of a dark amber tint

Tests —Pilocarpine Salicylate melts at about 120° C (248° F) and dissolves readily in Water, forming a solution which is faintly acid in reaction towards blue Litmus paper It dissolves in concentrated Sulphuric Acid without change of colour, but in Fuming Nitric Acid it forms a yellowish-brown solution The aqueous solution should yield a whitish amorphous precipitate with Potassiummercuric Iodide (Mayer's) Solution, and with Iodo-potassium Iodide (Wagner's)

reagent a brownish precipitate. The addition of Sodium Hydroxide Solution to a concentrated aqueous solution of the salt causes a whitish turbidity setting out into only drops soon dissolving in an excess of the Hydroxide Solution. The addition of Ammonia Solution to the concentrated aqueous solution of the salt should cause no precipitate, when dissolved in Water it yields on the addition of Ferric Chloride T.S. a deep violet coloration, the Salicylic Acid separated from the salt should possess the m.p. and answer the tests characteristic of Salicylic Acid given under that heading. When ignited with free access of air it should leave no weighable residue.

PILULÆ

PILLS

FR, PILULES, GER, PILLIN, ITAL, PILLORI, SPAN, PILDORAS

This class of medicine, so convenient and portable, was introduced in the earliest Pharmacopœias, and some of the formulas remain almost unchanged. The *Pilula Rufi* (*Pilula Aloes et Myrrhæ*) has for at least two hundred years maintained practically the same composition, but in *B.P.* '98 the Saffron is omitted.

Excipients for pills are of two kinds (1) those which are more or less fluid, and employed to bind together powders, or to impart the necessary moisture to adhesive substances, (2) those, generally in powder, which are intended to absorb moisture and give solidity to the mass. Of the former, 'Dispensing Syrup' (equal volumes of Alcohol (90 p.c.), Glycerin, Syrup and Mucilage) and 'Diluted Glucose' (Glucose 3, Syrup 1), are most in request, Alcohol (60 p.c.) also is very useful. Glycerin by itself is distinctly inferior to the foregoing. Glycerin of Tragacanth is much employed, but in the majority of cases where it would be used Glucose or 'Diluted Glucose' is preferable. Of the powders, that of Liquorice root is most useful when moisture is to be absorbed and no binding power is required. An unexpected exception is the case of Carbolic Acid, which makes a very good plastic mass with twice its weight of Liquorice powder (when well worked together the result is very satisfactory). When more plasticity is required the absorbent powder is supplemented by Compound Tragacanth Powder or powdered Gum Acacia. For essential Oils this condition is best obtained by the use of powdered Curd Soap, as a rule, 1 minim of the Oil will require half a grain of the Soap and 2 grains of the Liquorice. A good powder to mix with small doses of powerful medicines is the '**Diluting Mixture**' (Sugar of Milk 3, and Compound Tragacanth Powder 1), which will make a good pill with 'Diluted Glucose' q.s. A mixture of Paraffins (*Massa Paraffini*), or with Kaolin (*Massa Kaolini*), is used for substances which are readily reduced by organic matter, such as the Permanganates and the salts of Gold and Silver. It 'goes without saying' that an excipient must not be chemically incompatible with the other ingredients, but there is not much opportunity for such an occurrence with those above selected.

Coatings—Pills have been finished in various ways rolled in Flour, Starch, Magnesia, Liquorice powder, and in Lycopodium, or a

mixture of these, enveloped in Silver or Gold Leaf; coated with Ether-alcoholic solution of Tolu, or better with **Sandarach Varnish** (Ether 2, Absolute Alcohol 6, Sandarach 3), or with Gelatin or French Chalk. A good mucilage for the white coating to pills is Powdered Tragacanth, 1, Powdered Gum Acacia, 4, Diluted Acetic Acid, 8, Distilled Water, 40, or it can be made with Chloroform Water in place of Distilled Water, omitting the Acetic Acid. Another protective coating is **Salol Varnish** (Salol 1, Sandarach Varnish 5). Pills containing substances exceedingly soluble in Alcohol should not be varnished, as the varnish may dissolve some part of the pill.

When pills are intended to pass through the stomach, and to be disintegrated in the intestine, they are coated with a solution of Keratin, *see* p 710

PIMENTA.

PIMENTO.

FR, PIMENT DE LA JAMAIQUE, GER, ENGLISCHES GEWURZ, ITAL, PIMENTI, SPAN, PIMIENTA DE LA JAMAICA

The dried, full-grown, unripe Fruit of *Pimenta officinalis*, Lindley.
From the West Indies

Medicinal Properties.—A warm, aromatic stimulant and carminative, like Cloves, used as an adjuvant to tonics and purgatives

Dose.—10 to 30 grains = 0.65 to 2 grammes, in powder

Prescribing Notes.—The Oil may be given on sugar, or in pill with 1 part of P. to 10 parts of Sugar, *see* p 897

Official Preparations.—Aqua Pimentæ and Oleum Pimentæ

Foreign Pharmacopœias.—Official in Mex (Pimienta Gordal), Port (Pimienta da Jamaica), Span (Pimienta de la Jamaica), US Not in the others

Descriptive Notes.—The official fruit is known in commerce as Allspice or Jamaica Pepper, its resemblance in shape to Pepper is also indicated by the German name, Clove Pepper (Nelkenpfeffer). Pimenta being the Spanish for Pepper, the corrupted name Pimento was applied to it in the West Indies, and the same application of the name is made in France, where Allspice is termed Piment des Anglais, and the Capsicum or Guinea Pepper, Piment des Jardins.

The fruit is dried before it ripens, since it loses much of its essential Oil when ripe. It varies in size from about $\frac{1}{10}$ to $\frac{3}{10}$ in (2.5 to 7.5 mm) in diameter ($\frac{1}{2}$ to $\frac{1}{3}$ in, 5 to 8 mm, BP, 5 to 7 mm, USP). The remains of a four-toothed calyx crown the apex of the fruit, which is two-celled, each cell containing a reniform seed with a large, spirally-coiled embryo. Both pericarp and seed contain oil cells, but they are most numerous in the former. Pimento is produced by *Pimenta officinalis*, Lindl. The fruit of the allied *P. acris*, Wright (*Myrcia acris*, DC) is similar in appearance and sometimes occurs

in commerce, but has five calyx teeth. The leaves of this species are used in the manufacture of Bay Rum, but have a flavour different from Allspice, the leaves are known in commerce as 'Bay Leaves'. A better term would be West Indian Bay Leaves, since the term Bay Leaves properly pertains to those of *Laurus nobilis*, L. The fruits of Tobago or Mexican Allspice (*Eugenia Tabasco*, G. Don) are liable to be mistaken for the official article. They are but rarely met with, but are larger, paler brown and less aromatic.

The distinctive microscopical characters of the powdered fruit are the cluster crystals and rhomboidal crystals of Calcium Oxalate, small, thick-walled cells containing Resin, short, thick-walled, simple, tapering hairs, sclerenchymatous cells with branching pores, compound small starch grains, and spherical oil cells.

Powdered Pimento is stated by Moller (*Lehrb Pharmacognosie*, p 144) to have been adulterated with Pear stalks and Clove stalks, and he gives the methods for detecting these adulterations.

Tests—Pimento yields from 3 to 4 p.c. of ash, and should not exceed 5 p.c. It contains 3 to 4½ p.c. of volatile Oil and some quantity of Tannin.

Preparations

AQUA PIMENTÆ —PIMENTO WATER

Pimento, bruised, 4, Water, 160, distil one half (1 in 20)
Now 1 in 20 instead of 1 in 11½

Dose—1 to 2 fl. oz. = 28.4 to 56.8 c.c.

OLEUM PIMENTÆ —OIL OF PIMENTO

A pale yellow, or yellowish brown, oily liquid, heavier than Water, having a pleasant, clove-like odour and pungent spicy taste, distilled from Pimento. Yield about 3 to 4½ p.c.

On exposure to the air the colour darkens and the oil becomes thicker. It should therefore be kept in well-stoppered bottles of a dark amber tint and protected from the light. It contains a large percentage of Eugenol and a sesquiterpene.

Solubility.—In all proportions of Alcohol (90 p.c.), about 1 in 50 of Alcohol (60 p.c.)

Dose—½ to 3 minims = 0.03 to 0.18 c.c.

Not Official—Spiritus Myrciæ, Spiritus Pimentæ

Foreign Pharmacopœias—Official in U.S. Not in the others.

Tests—Oil of Pimento has a sp. gr. of 1.030 to 1.050. The B.P. states not below 1.040, the U.S.P. from 1.033 to 1.048 at 25° C (77° F). It is officially required to form a semi-solid mass when shaken with an equal volume of strong Ammonia Solution. The U.S.P. requires that it shall form a semi-solid mass when mixed with an equal volume of concentrated Sodium Hydroxide Solution, that it shall be miscible in all proportions with Alcohol (90 p.c.) and also soluble 1 in 2 of Alcohol (70 p.c.). 1 minim dissolved in 60 minims of Alcohol (90 p.c.) and treated with 1 minim of very dilute Ferric Chloride Solution assumes a fine indigo colour. Oil of Cloves, which Oil of Pimento very much resembles in chemical constitution, also

conforms to this test. It has been recommended that the Oil be required to indicate 65 p.c. of Eugenol by the Potassium Hydroxide method. The *U.S.P.* Oil is required to contain not less than 65 p.c. by volume of Eugenol as determined by measuring the portion undissolved when a measured quantity of 10 c.c. of the Oil is shaken for 5 minutes with 100 c.c. of Potassium Hydroxide T.S., the liquids being allowed to separate, sufficient Potassium Hydroxide T.S. added to raise the lower limit of the oily layer to the zero mark on the scale, a volume of not more than 3.5 c.c. should remain unabsorbed.

Not Official

SPIRITUS MYRCIÆ Bay Rhum—Oil of Myrcia, 0.8, Oil of Orange Peel, 0.05, Oil of Pimenta, 0.05, Alcohol (95 p.c.), 61, Water, *q.s.* to make 100—*U.S.P.* 1890

Spiritus Pimentæ *Syn.* Spiritus Myricæ Bay Rhum—Oil of Pimento Leaves, 0.75, Oil of Orange Peel, 0.05, Oil of Pimento, 0.05, Alcohol (90 p.c.), 64, Distilled Water, *q.s.* to produce 100—*B.P.C.*

PINI OLEUM.

OIL OF PINE

Colourless, or pale yellow, limpid oily liquid possessing an agreeable characteristic pine odour. It is distilled from the fresh leaves of *Pinus Pumilio*, Haenke.

It should be kept in well-closed glass bottles preferably of a dark amber tint.

Pine Oil contains Pinene, Lævo-phellandrene, Sylvestrene, Bornyl Acetate, Dipentene, and Cadinene.

'Pinol' and 'Pumiline' are commercial varieties of this Oil.

Solubility.—About $\frac{2}{3}$ dissolves 1 in 5 of Alcohol (90 p.c.), but the remaining $\frac{1}{3}$ is much less soluble.

Medicinal Properties.—The vapour or spray is a stimulating disinfectant expectorant in chronic catarrhal affections of the respiratory passages. The Oil is applied externally in rheumatism. Internally, as a disinfecting expectorant, the dose is 1 to 5 minims taken on sugar, or in the form of jujube or pastil.

Dose.—1 to 5 minims = 0.06 to 0.3 c.c.

Not Official—Extractum Pin. Pumilionis, Linctus Pini (Cypin et Hierom, Ruxir Pini et Terpin et Acetomorphinæ, Sympus Pin. Pumilionis and Vapor Olei Pin.

Foreign Pharmacopœias—Official in Austr. and Swiss. Not in the others.

Tests.—Pine Oil has a sp. gr. of from 0.865 to 0.875, the *B.P.* gives 0.865 to 0.870. It is lævogyrate, the optical rotation being from -5° to -9° ; the *B.P.* states -5° to -10° at 15.5° C. (60° F.). The *B.P.* requires that the *B.P.* should be below 160° C. (320° F.).

Not Official.

EXTRACTUM PINI PUMILIONIS—A liquid extract, of a brown colour, prepared from the young shoots of the *Pinus Pumilio*. It is soluble in Water, and is used in baths.

LINCTUS PINI TERPIN ET HEROIN *Syn* Elixir of Pine Terpene and Heroin—Pine Oil, 1 fl oz, Alcohol (90 p c), 5 fl oz, Terpene Hydrate, 40 grains, Glycerin, 5 fl oz, Light Magnesium Carbonate, 3 oz Saffron Tincture, 5 dm, Heroin Hydrochloride, 3½ grains, Syrup, q s to make 20 fl oz —*Martindale*

Dissolve the Terpene Hydrate in the Alcohol and Heroin Hydrochloride in the Syrup, and proceed as in preparing Syrupus Pini Pumilionis.

Elixir Pini et Terpin et Acetomorphinæ *Syn* Linctus Pini et Terpin et Acetomorphinæ—Oil of Pine, 5, Terpin Hydrate, 0.50, Acetomorphinæ Hydrochloride, 0.05, Alcohol, 25, Tincture of Saffron, 3, Glycerin, 25, Light Magnesium Carbonate, 15, Syrup, q s to produce 100 —*B P C*

SYRUPUS PINI PUMILIONIS—Pine Oil, 1 oz, Alcohol (90 p c), 5 oz, Saffron Tincture, 5 dm, Glycerin, 5 oz, Syrup, q s to make 20 fl oz. Rub the Pine Oil with 3 oz Light Magnesium Carbonate, then add the Alcohol, Glycerin, and Syrup, in parts, filter. Dose—1 dm (3.5 cc) —*Martindale*

This has been incorporated in the *B P C*

VAPOR OLEI PINI—Oil of Pine, 10, Magnesium Carbonate (light), 5, Distilled Water, q s to produce 100 —*B P C*

A similar inhalation appears below under the title Vapor Olei Pini Sylvestris

Not Official.

PINI SYLVESTRIS OLEUM

Under this name several varieties of Pine-needle Oil are supplied.

A colourless, or nearly colourless, limpid oily liquid with an agreeable odour, distilled from the fresh Leaves of various species of *Pinus*. The Oil distilled from the leaves of *Pinus sylvestris*, L., is not now obtainable in commerce.

Solubility—1 in 5 to 10 of Alcohol (90 p c), depending on the variety, in all proportions of Absolute Alcohol.

Medicinal Properties—Similar to those of Oil of Turpentine. It is also used externally in rheumatism, and as an inhalation or spray with hot Water in chronic laryngitis, bronchitis and phthisis.

Foreign Pharmacopœias—Official in Hung, sp gr 0.872, Russ (Oleum Pini Foliorum), sp gr 0.870 to 0.880. Not in the others.

Tests—The Oil should have a sp gr of not below 0.880. Rotation varies with the time of year at which the Oil is collected, climate and locality. Not more than 15 p c should distil below 170° C (338° F). Many Oils sold as *Pinus Sylvestris* yield on fractionation 60 to 70 p c, boiling below 167° C (332.6° F).

VAPOR OLEI PINI SYLVESTRIS—Oil of Scotch Pine, 40 minims; Light Magnesium Carbonate, 20 grains, Water, q s to produce 1 fl oz —*Throat*,

1 fl, dm, in 20 fl oz of Water at 140° F for each inhalation,

PIPER NIGRUM.

BLACK PEPPER

FR, POIVRE NOIR, GER, SCHWARZER PFEFFER, ITAL, PEPE NERO, SPAN, PIMIENTA NEGRA

The dried unripe Fruit of *Piper nigrum*, L

The ash of genuine Black Pepper varies from 4 to 6 p.c.
Chiefly from the East Indies

Medicinal Properties.—Carminative and stomachic Chiefly used to assist gastric digestion and relieve colic and flatulence Useful in hæmorrhoids and in urethritis

Official Preparation—Confectio Piperis Contained in Pulvis Opi Compositus

Not Official—Oleoresina Piperis, Piperina, Piperidine, Piperidino Guaiacolate, Piperidine Acid Tartrate

Foreign Pharmacopœias—Official in Austr, Belg, Jap, Mex (Pimenta Negra), Port (Pimenta), U S Not in the others

Descriptive Notes—Black Pepper consists of the dried unripe fruits of *Piper nigrum*. The fruits are black, nearly spherical, about $\frac{1}{2}$ in (5 mm) in diameter, wrinkled, slightly pointed below from the remains of a very short pedicel, and showing traces at the apex of a 3 to 4 lobed stigma. The single seed fills the fruit and contains a small cavity at the apex where the embryo should be developed in the ripe fruit. The albumen is horny externally and starchy inside. The taste is pungent and the flavour and odour characteristic. It is largely imported from Singapore, Malabar, (Tellicherry and Aleppy), Ceylon, Siam, and Mangalore. The last named is large and has a flavour resembling Bay Leaves. Penang Pepper is preferred for its strength and Sumatra Pepper (Achou and Lampong) for its colour. The heavy or shot Pepper of Tellicherry and Ceylon is preferred for grinding. Black Pepper is collected as soon as the lowest fruits on the spikes turn red, since it loses some of its pungency as it ripens, although it improves in flavour. White Pepper consists of the ripe fruit with the pericarp removed by soaking in water and rubbing. The Tellicherry and Ceylon kinds of White Pepper are considered to be the best. The Black Pepper of the B P is described as $\frac{1}{2}$ in (5 mm) in diameter and almost black in colour, that of the U S P should be greyish-black, and 4 to 5 mm in diameter.

The distinctive microscopical characteristics of Black Pepper are the parenchymatous cells with brown contents more or less surrounded by thin-walled parenchymatous cells, and the sclerenchymatous cells of the inner layer of the pericarp, which have larger cavities that are clear, and unequally thickened walls, the small starch grains (0.002 mm, U.S.P.), and the oil cells.

Whole Pepper is rarely adulterated. Ground Pepper has been adulterated with rice, which has a distinctive starch; ground Olive stones, which become pulverulent when boiled with solution of ammonia in acetic acid; and, according to Müller, with ground Mustard, Capsicum,

and other farinaceous substances, some illustrations of which are given in his *Lehrbuch der Pharm* 1889, pp 136-7

Tests—Black Pepper leaves on ignition from 4 to 6 p c of ash, and should not leave more than 7 p c The *USP* ash limit is 7 p c.

It has been recommended that the characters of the powder should be given together with an ash and possibly an Oleo-resin standard The matter soluble in hot Alcohol amounts to about 7 p c.

Preparation

CONFECTIO PIPERIS CONFECTION OF PEPPER

Black Pepper, 2, Caraway fruit, 3, Clarified Honey (by weight),
15 (1 in 10)

Dose —60 to 120 grains = 4 to 8 grammes

Not Official

OLEORESINA PIPERIS (US)—Obtained from Pepper by exhaustion with Acetone, evaporation of the solution, and finally separation of the crystals of Piperine by straining the Oleo resin through Cotton-Wool

Average Dose — $\frac{1}{2}$ grain = 0.032 gramme

This has been incorporated in the *BPC*

PIPERINA Piperine $C_{17}H_{19}NO_3$, eq 283.05—It is described in the *USP*, as a feebly basic substance obtained from Pepper and other plants of the Piperaceæ Allen describes it as an alkaloid existing in various plants belonging to the Piperaceæ, and as the characteristic principle of both black and long Pepper. It forms colourless or pale yellow four-sided monoclinic prisms, odourless, and at first tasteless, but subsequently developing a sharp biting taste Insoluble in cold Water, very slightly soluble in boiling Water, is sparingly soluble in Ether, but dissolves readily in Chloroform and Benzene

It possesses antipyretic properties, but it is not the active principle of Pepper

Official in Mex

Average Dose —3 grains = 0.2 gramme

Tests—Piperine melts at 128° to 129° C (262.4° to 264.2° F) The *USP*, states at 130° C (266° F) It dissolves in Alcohol, the alcoholic solution being neutral in reaction towards Litmus paper and optically inactive It dissolves in Sulphuric Acid with the formation of an orange red coloration, becoming brown on warming or standing and disappearing on dilution with Water On the addition of Nitric Acid it acquires an orange red coloration, which is turned to a blood-red colour by the addition of an excess of Potassium Hydroxide Solution Sulphuric Acid containing about half its volume of Formaldehyde Solution produces a permanent green solution When mixed with Sulphuric Acid containing a crystal of Potassium Bichromate a purple coloration is immediately developed, changing on stirring to a reddish-brown solution, which becomes greenish on the addition of Water On the addition of Iodo potassium Iodide Solution to a hot alcoholic solution acidified with Hydrochloric Acid an Iodo compound is formed, which separates on cooling into fine steel blue needles When boiled with Potassium or Sodium Hydroxide Solution, Piperine undergoes saponification, being converted into Sodium Piperinate and Piperidine Piperinic Acid melts at about 215° C (419° F) When ignited with free access of air Piperine should leave no weighable residue

PIPERIDINE ($C_4H_{11}N$, eq 84.49)—A colourless limpid liquid Possessing a strongly alkaline reaction and a strong odour resembling Pepper It is a powerful base, produced by the hydrolysis of Pipefine, the alkaloid occurring in Pepper, or synthetically by the reduction of Pyridine by nascent Hydrogen.

Tests—It boils at 106°C (222.8°F), and distils unchanged at that temperature. It dissolves in Alcohol, the solution possessing a strongly alkaline reaction to litmus. It is determined by titration with Normal or Tenth-normal Vol. or Sulphuric Acid Solution, using either Litmus or Methyl Orange Solution as an indicator of neutrality. 1 cc of Normal Volumetric Acid Solution is equivalent to 0.08449 gramme of Piperidine. It rapidly absorbs Carbon Dioxide from the air.

PIPERIDINE GUAIACOLATE (Guaiaperol)—A compound of Piperidine and Guaiacol. A yellowish-white crystalline body, having a faint odour of Guaiacol. Soluble in Water. Mineral acids and alkalis decompose it into its constituents. Recommended in phthisis.—*B M J* '97, 1 136, *J.C.S. Trans* '98, 145.

Dose—5 to 30 grains = 0.32 to 2 grammes

PIPERIDINE ACID TARTRATE—A white crystalline powder possessing a faint odour. Readily soluble in Water. Introduced as a solvent for gouty deposits, Uric Acid gravel and calculi. It increases the solvent power of serum for Sodium Bicarbonate to a much larger extent than Piperazine, Lysidine or Urotropine.—*L* '98, 1 198, 280, 345, 438, 507.

Dose.—10 to 15 grains = 0.65 to 1 gramme

Calculusol is stated to be a mixture of Piperidine Para-Sulphamine Benzoate and Potassium Bicarbonate.

Not Official.

PIPERAZINE.

HEXAHYDROPIRAZINE

$(\text{C}_2\text{H}_4\text{NH})_2$, eq 85.52.

Colourless deliquescent crystals, readily soluble in Water.

Piperazine (Diethylene-diamine) = $\text{C}_4\text{H}_{10}\text{N}_2$. Ammonia on Ethylene Bromide or Chloride. Or by Zinc dust and metallic Sodium.

Medicinal Properties—It has in the laboratory a powerful solvent action on Uric Acid, the Piperazine Urate being about seven times more soluble than Lithium Urate, but whether it has a similar action in the body is doubtful. Recommended for gouty rheumatoid affections and renal calculus and colic.—*T G* '93, 19, '94, 1 1291. *B M J* '93, 1 20, 1 184, 1 265.

Little or no effect in gouty states.—(Sir Wm Roberts and *Fr* '93, 1 50; in diabetes.—*B M J E* '93, 1 72, action as a Uric Acid solvent.—*L* '96, 1 1901.

Piperazine did not delay the conversion of gelatinous Sodium Bicarbonate into the crystalline variety, and the conversion when once started was but slightly slowed by the presence of this substance.—*B M J* '00, 1 836, *L* '00, 1 931. 0.05 gramme dissolved in 0.5 cc Water, injected in gouty tophi.—*B M J* '99, 1 56.

Dose—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—Usually given in mixture, as a granular effervescent preparation, containing 2½ grains in each 60 grains.

Foreign Pharmacopœias.—Official in *Fr*.

Tests.—Piperazine melts when anhydrous at 104° to 107°C (219.2° to 224.6°F); *Fr. Code* gives 104°C (219.2°F). It boils at 145°C (293°F), *Fr. Code* gives between 145° and 146°C (293° and 294.8°F). It mixes readily in Water, the aqueous solution having a strongly alkaline reaction towards red litmus paper. On the addition of Potassium Bismuth Iodide Solution, to a slightly acid solution, a white precipitate is thrown down. On the addition of Picric Acid Solution, a yellow precipitate is thrown down, exhibiting a

characteristic microscopical appearance is thrown down. A glass rod moistened with Hydrochloric Acid gives off dense white clouds when held over a crystal of Piperazine. The aqueous solution affords with Potassio mercuric Iodide (Mayer's) Solution a white precipitate, with Mercuric Chloride Solution a white precipitate, with Tannic Acid Solution an amorphous precipitate soluble in hot Water. The aqueous solution when acidified with Hydrochloric Acid yields with Platinic Chloride Solution a reddish yellow precipitate, and if the solution be not too dilute Gold Chloride Solution throws down a crystalline double salt soluble in hot Water. Piperazine is not altered by aqueous Chromic Acid Solution, but Potassium Permanganate oxidises it quickly in the cold. Piperazine absorbs Carbonic Acid rapidly from the air, being converted into the Carbonate which melts at 162° to 165° C (323° F to 329° F). It should yield no brown or reddish-brown coloration with Alkaline Potassio mercuric Iodide (Nessler's) Solution, indicating the absence of Ammonium salts. When acidified with Nitric Acid the aqueous solution should yield neither a turbidity nor precipitate with either Silver Nitrate Solution or Barium Nitrate Solution, indicating the absence of Chlorides and Sulphates. When strongly heated it should completely sublime without leaving a weighable residue indicating the absence of mineral impurities.

LYCETOL (Dimethylpiperazine Tartrate)—A white powder readily soluble in Water, possessing an acid taste. Recommended in chronic gout and rheumatism.

Dose—5 to 10 grains = 0.32 to 0.65 gramme

LYSIDINE (Ethylene-ethenyl diamine)—A reddish white, crystalline substance, very hygroscopic, with a peculiar odour. Commercially it is sold in the form of a 50 p.c. solution.

A diuretic, recommended in gout and as a solvent of Uric Acid deposits.—*B M J* '96, 11 901

It has an influence in increasing the solvent power of serum for Sodium Bicurate and of urine for uratic deposit.—*L* '98, 11 203

Though it delayed the conversion of the gelatinous Sodium Bicurate into the crystalline form, yet when the conversion was once started it had practically no effect in slowing it.—*L* '00, 1 931, *B M J* '00, 1 836

Dose (of the liquid)—30 to 60 minims = 1.8 to 3.6 c.c., well diluted with Water or aerated Water.

Lysidine Acid Tartrate—A white powder soluble in Water.

Dose—15 to 30 grains = 1 to 2 grammes

Piperazine Quinate or Kinate (Sidonal)—A white granular powder, readily soluble in Water, the solution having a pleasant slightly acid taste. Recommended as a solvent of Uric Acid and gouty deposits.—*B M J* '01, 1 1408

Dose—5 to 15 grains = 0.32 to 1 gramme

New Sidonal (Quinic Anhydride)—A white, crystalline powder, readily soluble in Water. Introduced for the treatment of gout.

Dose—5 to 15 grains = 0.32 to 1 gramme

Not Official PISCIDIA

Syn—JAMAICA DOGWOOD

The Root-bark of *Piscidia erythrina*, Lam.

The shrub is a native of South America and the West Indies, where it has been used for stupefying fish.

Medicinal Properties—A sedative in nervous irritability and in irritant cough, an antispasmodic in asthma.

Has been used in neuralgia and toothache.—*P J* (3) xvi 1014

Official in Mex (Colorin de peces).

EXTRACTUM PISCIDIÆ LIQUIDUM—1 fl oz is equal to 1 oz of the
100t

Dose—30 to 120 minims = 1 8 to 7 1 c c.

Extractum Piscidiæ—The above evaporated to an extract for pills
Dose, 1 to 5 grains = 0 06 to 0 32 gramme

PIX BURGUNDICA.

BURGUNDY PITCH

FR., POIX DE BOURGOGNE, GER, FICHTENHARZ, ITAL, PECE DI BORGOGNA;
SPAN, PEZ DE BORGONA

An opaque or translucent yellowish- or reddish-brown brittle solid, possessing a terebinthinate odour Imported from Germany The prepared resinous exudation obtained from the stem of *Pinca excelsa*, Link

It is the Thus of Frankincense of Lond and Dub
exudes from the spruce fir, and when melted and strained
Pitch

Solubility.—Almost entirely dissolves 1 in 20 of Alcohol (90 p c), the greater part dissolves 1 in 1½ of Glacial Acetic Acid

Medicinal Properties.—The plaster is applied to the chest in chronic pulmonary complaints, to the loins in lumbago, to the joints in chronic articular affections, and to other parts to relieve local pain of a rheumatic character It acts as a counter-irritant

Official Preparation—**Emplastrum Picis**

Foreign Pharmacopœias—Official in Belg (Pix de Bourgogne), Hung (Resina Pini Burgundica), Ital (Poix de Bourgogne), Mex and Span (Pez de Borgona), Port (Pez de Borgonha), Swiss (Resina Pini) Not in Austr, Dan, Dutch, Ger, Jap, Norw or Russ

Descriptive Notes.—True Burgundy Pitch is rarely met with in English commerce, its place being taken by a factitious article made with ordinary Resin, Turpentine and Palm Oil, and sold in bladders The genuine article is produced in Finland and the Black Forest and, as described by Hanbury, has an agreeable resinous odour, but when old it recalls that of Castor Oil It is a yellowish-brown, brittle and hard, but gradually becomes more so the vessel in which it is kept, has a clean and shining surface It gives off an aromatic odour when heated, it does not exhibit a crystalline structure The fat present in the factitious article will not dissolve in Glacial Acetic Acid, the taste should be sweet, aromatic and without bitterness

Tests.—Burgundy Pitch is officially required to be readily soluble in Glacial Acetic Acid

Preparation

EMPLASTRUM PICIS.—**PITCH PLASTER**

Burgundy Pitch, 26, Frankincense, 13, Resin, 4½, Yellow Beeswax, 4½, Olive Oil (by weight), 2, Distilled Water, 2; melt and evaporate to the consistence of a plaster

The Expressed Oil of Nutmeg is omitted in B P 1898.

Foreign Pharmacopœias—Official in Fr, Port and Span Yellow Wax 1, Burgundy Pitch 3, Ital (*Empiastro Adesivo*), Yellow Wax 3, Burgundy Pitch 7, Diachylon Plaster 40, Mex (*Emplastro Aglutinante*), Pitch 74, Elemi 10, Sesame Oil 6, Yellow Wax 10 Not in the others

PIX CARBONIS PRÆPARATA.

PREPARED COAL TAR

Prepared from commercial Coal Tar by dissipating all constituents volatile below 120° F (48 9° C), by keeping it at that temperature for 1 hour in a shallow vessel

The solution is used in chronic eczema as a Lotion, 1 to 20 or more of Water, or as an Ointment 1 to 8 It is frequently prescribed with the Liquor or the Glycerole of Lead

Official Preparation—Liquor Picis Carbonis

Not Official—Liquor Carbonis Detergens, Lotio Plumbi et Picis, Plâtre coaltaré (Vet), Unguentum Picis Carbonis, Unguentum Picis Carbonis Compositum, Unguentum Petrolati Compositum

Foreign Pharmacopœias—Official in Dutch and Swiss (*Pix Lithanthracis*), Fr (*Goudion de Houille*) Not in the others

Preparation

LIQUOR PICIS CARBONIS—SOLUTION OF COAL TAR

Digest for 2 days at 120° F (48 9° C) 1 (by weight) of Prepared Coal Tar in 5 of a Tincture of Quillaja (1 in 10, Alcohol 90 p c), decant or filter when cold (1 in 5)

Foreign Pharmacopœias—Liquor Picis Lithanthracis —*Pix Lithanthracis* 1, Alcohol (90 p c) 1—*Dutch*

Not Official

LIQUOR CARBONIS DETERGENS—An Alcoholic solution of Coal Tar It is almost black, smells strongly of Naphthalene, and is of light sp gr Used externally in chronic scaly skin diseases diluted about 1 in 20 of Water

Coal Tar in dermatological practice —*B M J E* '94, 11 88

LOTIO PLUMBI ET PICIS—Strong solution of Lead Acetate, 10 minims, Solution of Coal Tar, 20 minims, Water, to 1 fl oz —*London*

PLÂTRE COALTARÉ (Vet) —Coal Tar, 1, Calcium Sulphate (Moulder's Plaster), 20 —*Fr*

UNGUENTUM PICIS CARBONIS—Solution of Coal Tar, $\frac{1}{2}$ fl drm, Soft Paraffin, yellow, to 1 oz —*St Thomas's*

Solution of Coal Tar, by weight, 6, Soft Paraffin, yellow, q s to produce 100 —*B P C*

UNGUENTUM PICIS CARBONIS COMPOSITUM—Solution of Coal Tar, $\frac{1}{2}$ fl drm, Ammoniated Mercury, 15 grains, Soft Paraffin, yellow, to 1 oz, —*St Thomas's*

Solution of Coal Tar, by weight, 6, Ammoniated Mercury, 3, Soft Paraffin, yellow, q s to produce 100 —*B P C*

Unguentum Petrolati Compositum—Solution of Coal Tar, $\frac{1}{2}$ drm, Ammoniated Mercury, 10 grains, Soft Paraffin, to 1 oz —*St John's*

PIX LIQUIDA.

TAR.

FR, Goudron Végétal, GER, Holztheer, ITAL, Catrame Vegetale;
SPAN, Brea

A thick, dark brown or brownish-black, bituminous fluid or semi-fluid, having a strong, peculiar, empyreumatic terebinthinate odour. Obtained by destructive distillation from the wood of *Pinus sylvestris* and other species of *Pinus*, *USP* says *Pinus palustris*. Known commercially as Stockholm Tar.

Wood Tar contains Guaiacol and Cresol. Coal Tar yields Phenol and Cresol.

Solubility.—In less than its own bulk of Alcohol (90 p c) or Chloroform and separates on the addition of Water, soluble 1 in 3 of Solution of Sodium Hydroxide (4 p c), slightly soluble in Olive Oil or Oil of Turpentine.

Medicinal Properties.—Similar to Turpentine. May be used as a disinfectant expectorant in chronic bronchitis and winter cough, taken internally or inhaled from steaming Water. The ointment is used in leprosy, pruritus, and also for some chronic skin diseases, such as eczema and psoriasis.

Dose.—5 to 10 minims = 0.3 to 0.6 c c, but the dose may be increased gradually.

Prescribing Notes.—Best given in capsules. Tar varies slightly in substance, and is very difficult to form into pills of a convenient size, and so much excipient, that a 5-grain pill will contain very little Tar. Liquorice Root and Lycopodium have been recommended for this purpose, neither of them alone yields a satisfactory mass. Equal weights of Tar, (1 lb) Soap, Powdered Liquorice Root, and Powdered Gum Acacia, make a good plastic pill, the quantity of Tar which can be worked up with this mixture will vary according to the consistence of the Tar.

Official Preparation.—Unguentum Picis Liquidæ

Not Official.—Tinctura Picis, Oleum Picis, Aqua Picis, Capsule Picis, Oleum Picis Liquidæ, Oleum Picis, Pigmentum Picis Liquidæ, Pilule Picis, Syrupus Picis Liquidæ, Syrupus Picis cum Codeno, Vasolimentum Picis, Parogenum Picis, Black Pitch.

Foreign Pharmacopœias.—Official in all, Dan, Norw and Swed (Pyroleum Pinii), Fr (Goudron végétal), obtained from *Pinus maritima*, Ital (Catrame vegetale), Mex (Alquitran), Port (Alcatrazo), Span (Brea), Russ has Pix Solida also.

Descriptive Notes.—Official Tar is commonly known in commerce as Wood Tar, Archangel or Stockholm Tar, to distinguish it from Coal Tar. It is obtained by destructive distillation of the stumps and roots, chiefly of *Pinus sylvestris*, L., and *Abies Siberica*, Ledeb., in Northern Europe. Some varieties exhibit colourless crystals of Pyrocatechin, to which it owes its occasional granular appearance. The Pix Liquida of the *USP* is derived from *Pinus palustris*, Miller, and other species.

Tests.—Tar is required by the *BP* to have a sp gr of 1.02 to 1.15, the *PG* and *USP* state that it is heavier than Water. When shaken with Water the aqueous solution acquires a light brown

colour and a sharp and empyreumatic taste. The aqueous liquid has an acid reaction towards blue Litmus paper and affords, with dilute Ferric Chloride T S, a red coloration. According to the *P G*, 20 c c of Water which has been shaken with Tar assumes a greenish brown coloration on the addition of 2 drops of Ferric Chloride T S. The *P G* states that a mixture of equal volumes of Lime Water in which Tar has been shaken and Lime Water assumes a dark brown colour. When ignited with free access of air it should leave no weighable residue.

Preparation

UNGUENTUM PICIS LIQUIDÆ —Tar Ointment

Tar (by weight), 5, Yellow Beeswax, 2

This ointment is too hard for use. A proper consistence is obtained by replacing half of the Yellow Beeswax with Almond Oil (see Ung. Picis Molle).

Foreign Pharmacopœias —Official in Dan, Pitch 9, Lard 6 Potassium Carbonate 3, Water 2, Dutch (Ung. Picis), Pix Solida 3, Colophonium 4, Yellow Wax 2, Sesame Oil 12, also Ung. Picis Co., Fr (Pommade de Goudron), and Port, Tar 1, Lard 9, Jap, Wood Tar 10, Yellow Wax 4, Span, Tar 3, Lard 17, U S, Tar 50, Yellow Wax 15, Lard 35. Not in the others.

Not Official

UNGUENTUM PICIS MOLLE —Tar (by weight), 5, Yellow Beeswax, 1, Almond Oil, 1, melt together and stir till cold.

AQUA PICIS (Tar Water, Aqua Pyrolii Pini, Aqua di Catrame, Eau de Goudron) —Tar, 1, finely powdered, washed and dried Pumice, 3, Distilled Water, 200, agitate for 15 minutes, and filter.

Dose —From 1 to 2 pints daily, or may be used as a wash for ulcers and wounds.

The *B P C* adopts the strength given in *Ph. Ger*, Tar 1, Water 10.

Foreign Pharmacopœias —Official in Dutch, Tar 1, Pumice 3, Water 20. Fr (Eau de Goudron), Tar 1, Calcined Sand 3, Water 200, Ger, Jap and Swiss, Tar 1, Pumice 3, Water 10, Norw (Aqua Pyrolei Pini), 1 in 10, Mex (Aqua de Alqutran), Tar 5, Pumice 15, Water 1000, Port (Aqua de Alcatiao), and Ital (Acqua di Catrame), 1 in 40. Span (Aqua de Brea), 1 in 33, also (Solucion de Brea Alcalina), Bicarbonate of Soda 20, Tar 40, Distilled Water 85, Russ, Buch Tar 1, Water 90. Not in Austri, Hung, or U S.

CAPSULÆ PICIS —Capsules containing 5 minims = 0.3 c c

Dose —1 or 2 capsules

OLEUM PICIS LIQUIDÆ (Oil of Tar) —This volatile Oil, distilled from Tar, is official in U S as an almost colourless liquid when first distilled, but becoming dark reddish brown on keeping, sp gr about 0.892 at 25 °C (77 °F).

OLEUM PICIS RECTIFICATUM (Light Oil of Tar) —Colourless when first distilled, becoming brown on keeping, sp gr 0.840 to 0.870.

PIGMENTUM PICIS LIQUIDÆ —Tar, 1, Alcohol (90 p c), 1

Used as a stimulant in psoriasis and chronic dry eczema. Its use in eczema demands caution.

PILULÆ PICIS —Tar, Curd Soap, powdered Liquorice Root, and powdered Gum Acacia, equal weights mixed, and made into 5 grain pills.

Dose —2 or 3 pills thrice daily.

They are sometimes made of Black Pitch, and have been taken to relieve hæmorrhoids.

SYRUPUS PICIS LIQUIDÆ —Tar, 0.5, Alcohol (95 p c), 5, Magnesium Carbonate, 1, Sugar, 85, Water, q s to produce 100 —Mix the tar intimately in

a mortar with 1 of clean white Sand, add 10 of Water, and after loading the mass thoroughly with the pestle pour off the Water and throw it away. Treat the residue with the Alcohol, and when the Tar is dissolved add the Magnesium Carbonate and 5 of Sugar, and after thorough trituration add 40 of Water, stir occasionally during 2 hours and filter, dissolve the remainder of the Sugar in the clear filtrate by gentle heat, strain and add sufficient Water to make the product 100. *U. S. P.*

This has been incorporated in the *B. P. C.*

Dose—1 to 2 fl drms = 3.6 to 7.1 cc

Sometimes prescribed with Syrup of Wild Cherry Bark, and also with Codeine

Foreign Pharmacopœias—Official in *Fr.* (Simp de Goudron), Tar 10, Calcined and washed Sandstone 15, Distilled Water 1000, Sugar in of 18 to 10 of the liquid, *Span*, Solucion de Brea Alcalina 12, Tar ' ' ' ' Saccharum 64

SIRUPUS PICIS CUM CODEINO—Codeine 0.1, Proof Spirit *q s*, Syrup Picis Liquid to make 100. **Dose**—½ to 2 fl drms—*Swiss* 1893. Tar Water, 324, Sugar, 505, Glycerin, 150, Codeine, 1, Diluted Spirit (Alcohol 60 p c), 20—*Swiss* 1907

VASOLIMENTUM PICIS—Tar, 25, dissolved in Alcoholic Ammonia, 25, Simple Vasoliment, 75, mixed, evaporated on a water-bath to 100, and filtered—*Y. B. P.* 1901, 212, and *Hager*

Parogenum Picis *Syn* Tar Vasoliment—Tar, 25, Parogen, *q s* to produce 100—*B. P. C.*

BLACK PITCH.—There are three kinds, Archangel, Swedish, and that obtained from Gas Tar, the latter is without odour

Not Official

PLUMBUM

LEAD

Pb, eq 205.85

Lead occurs in nature as an Oxide, and as a Sulphide called *Galen*, saline combination, forming the native Lead Sulphate, Phosphate, Chromate, Molybdate, Tungstate, and Arsenate. The native Oxide is rare, but Galena, the ore from which nearly all the Lead of commerce is extracted is exceedingly abundant.

Fr., Plomb, *Ital.*, Piombi, *Mex.* and *Span.*, Plomo, *Port.*, Chumbo

Incompatibles

Antidotes

} Are given after Plumbi Subacetatis Liquor Fortis, p 919

Tests—Lead has a sp gr of 11.3, it fuses at 325° C (617° F). It is not affected in the cold by Sulphuric Acid, but when heated it dissolves with the evolution of Sulphur Dioxide. When heated in the air it is converted into Lead Oxide. It dissolves readily in diluted Nitric Acid, forming a solution which affords, with Sulphuric Acid, a white precipitate insoluble in Water, reprecipitating as the solution cools. It also reacts, treated so, with Hydrogen Sulphide, a black precipitate, insoluble in Hydrochloric Acid, insoluble in Potassium Hydroxide Solution and in solution of Ammonium Hydroxide, the black precipitate is decomposed by boiling with diluted Nitric Acid. Diluted Sulphuric Acid affords a white precipitate almost insoluble in Water and dilute Sulphuric Acid, insoluble in Alcohol (90 p c), but soluble in Ammonium Acetate Solution. Potassium Chromate Solution affords a yellow precipitate readily soluble in Potassium Hydroxide Solution, and in strong, hot Nitric Acid, sparingly soluble in diluted Nitric Acid, insoluble in Acetic Acid. Potassium Hydroxide Solution yields a white precipitate, soluble in excess of the reagent, insoluble in Ammonium Solution. A neutral solution affords, with

Potassium Iodide Solution, a yellow crystalline precipitate, soluble on boiling, and depositing again in brilliant golden crystals as the solution cools.

Lead salts are distinguished when in solution from those of any other metals by giving white precipitates with soluble Chlorides and Sulphates, insoluble in any dilute acid, yellow precipitates with Chromates and Iodides, a black precipitate with Hydrogen Sulphide from an acid solution. All these precipitates (except the Sulphides) are soluble in excess of hot Potassium or Sodium Hydroxide Solution.

PLUMBI ACETAS.

LEAD ACETATE

$\text{Pb}(\text{C}_2\text{H}_3\text{O}_2)_2, 3\text{H}_2\text{O}$, oq 376 15

FR, ACETATE NEUTRE DE PLOMB, GR, BLETACETAT, ITAL, ACETATO DI PIOMBO, SIAM, ACETATO PLUMBICO

Colourless, translucent prismatic crystals, or as masses of white monoclinic prisms possessing a faint odour of Acetic Acid and a sweet metallic and astringent taste.

It should be kept in well-closed bottles and in a cool atmosphere, as it is slightly efflorescent and liable to absorb Carbon Dioxide on exposure to the air.

Solubility—1 in 2 of Water, 6 in 1 of boiling Water, 1 in 20 of Alcohol (90 p c), 1 in 2 of Glycerin.

Medicinal Properties—In small doses it is sedative and astringent, lessening morbid mucous discharges and hæmorrhages in the gastro intestinal and genito-urinary tracts, and even diminishing natural secretions, hence it is useful in gastric ulcer, diarrhœa, dysentery, cholera, and in tubercular and typhoid ulceration. Used in phthisis to check excessive expectoration and to allay hæmorrhage, in bronchitis to abate profuse secretion. Its prolonged use requires caution, otherwise chronic Lead poisoning may be induced. It is often accompanied or followed by a small dose of Acetic Acid, as excess of Acid makes it less injurious to the system. Externally it is sedative, desiccant, and astringent, diminishing profuse discharges of ulcers, used for injection in gonorrhœa and other chronic inflammatory discharges.

Along with Opium, or better, as *Lotio Plumbi Evaporans cum Morphina*, it is a favourite application for sprains and bruises. The compound Lead suppositories are used for painful and bleeding piles.

A solution of this salt with a little sublimed Sulphur is stated (*B M J* '04, n 1749) to form an excellent test for *tinea versicolor*. A little of the lotion is applied to the part, and after the lapse of a short time, the *tinea* will appear clearly marked out in blackened patches.

In intestinal hæmorrhage 5 grains every hour.—*T G* '07, 324

Dose—1 to 5 grains = 0.06 to 0.32 gramme

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme

Prescribing Notes—May be given in pills with $\frac{1}{4}$ to $\frac{1}{2}$ of its weight of Compound Tragacanth Powder, and massing with 'Diluted Glucose' or Dispensing Syrup, q s in solution, with excess of Acetic Acid, with Opium in the official Pill, and Suppository.

Incompatibles—Sulphuric and Tannic Acids, and their salts, Chlorides, Iodides, and Phosphates

Official Preparations—*Pilula Plumbi cum Opio*, *Suppositoria Plumbi Composita*, and *Unguentum Plumbi Acetatis*. Used in the preparation of *Glycerinum Plumbi Subacetatis*, *Liquor Plumbi Subacetatis Fortis*

Not Official—*Lotio Plumbi Acetatis*

Antidotes—Same as under *Plumbi Subacetatis Liquor*

Foreign Pharmacopœias—Official in all, Austri, Ger and Swiss (*Plumbum Aceticum*), Hung and Russ (*Plumbum Aceticum Depuratum*), Dan, Dutch, Norw and Swed (*Acetas Plumbicus*), Plumbi Pl (Acetate Neutre de Plomb), Ital (*Acetato di Piombo*), Mex (*Acetato de Plomo*), Port (*Acetato de Chumbo*), Span (*Acetato Plumbico*)

Tests.—Lead Acetate loses its Water of crystallisation when heated to 40° C (104° F), suffering a loss of weight of 14.26 p.c. It dissolves in Water to form a clear solution, which should be only slightly acid in reaction towards blue Litmus paper, and which should only be slightly opalescent, the opalescence disappearing on the addition of a drop or two of Acetic Acid. The *USP* states that the aqueous solution has a neutral or slightly alkaline reaction. The *PG* states that the cold saturated aqueous solution of the salt has an alkaline reaction towards red Litmus paper, and on dilution a faintly acid reaction towards blue Litmus paper. The aqueous solution should answer the tests distinctive of Lead given under *Plumbum*. When mixed with Sulphuric Acid it evolves a strong acetous odour, when mixed with Sulphuric Acid and warmed with a small quantity of Alcohol (90 p.c) it evolves the distinctive odour of Ethyl Acetate. It is officially required to contain 99.8 p.c of pure crystallised Lead Acetate, as determined by precipitating 1 gramme of the salt dissolved in Water with Tenth-normal Volumetric Sulphuric Acid Solution. It should require for complete precipitation 53.1 c.c. The *USP* requires it to contain not less than 99.5 p.c of pure Lead Acetate, but does not give a method by which this percentage can be assured. The *PG* gives neither a requisite percentage of pure Acetate nor a method of determination. The use of Normal Volumetric Oxalic Acid Solution in the place of Normal Volumetric Sulphuric Acid Solution has been recommended (*PJ* '98, ii 531) as a more useful method of determining the amount of pure Lead Acetate present. The excess of Normal Volumetric Oxalic Acid Solution may be titrated by Tenth-normal Volumetric Potassium Permanganate Solution. This volumetric process has been adopted by the *USP* for the assay of the Lead Subacetate Solution.

The more generally occurring impurities are Copper, Iron, Zinc, Calcium, Magnesium, and salts of the alkali metals and Carbonates. The formation of a clear solution when the salt is dissolved in Water precludes the presence of Carbonates. The Potassium Ferrocyanide test described below indicates Iron and Copper if present. A test for the presence of Zinc, and an additional test for Iron, is afforded by Hydrogen Sulphide and Ammonia Water, the test is described below. Calcium, Magnesium, and the salts of the alkali metals, if present, may be detected in the filtrate from this test.

Potassium Ferrocyanide—Lead Acetate should give a clear, or at most only faintly opalescent, solution with 10 parts of Water (which has been recently boiled, *U S P*) and this solution should yield, with Potassium Ferrocyanide, a pure white precipitate, *P G* and *U S P*

Hydrogen Sulphide and Ammonia Water—If the Lead be precipitated from an aqueous solution of the salt, first by Hydrochloric Acid until precipitation ceases, then filtering and adding Hydrogen Sulphide and filtering again, a portion of the second filtrate should not be affected by the addition of a slight excess of Ammonia Water—*U S P*

Residue—Another portion of this second filtrate as above, when evaporated to dryness, should leave no residue—*U S P*

Preparations

PILULA PLUMBI CUM OPIO PILL OF LEAD WITH OPIUM

Lead Acetate, 12, Opium, 2, Syrup of Glucose, about 1

Dose—2 to 4 grains = 0.13 to 0.26 gramme

A 4 grain pill contains about 3 grains of Plumbi Acetas and $\frac{1}{2}$ grain of Pulvis Opii

Foreign Pharmacopœias—Official in Port, Lead Acetate, 5, Extract of Opium, 1, Extract of Liquorice, 14 Not in the others

SUPPOSITORIA PLUMBI COMPOSITA COMPOUND LEAD SUPPOSITORIES

3 grains of Lead Acetate, and 1 grain of Opium, in each with Oil of Theobroma

UNGUENTUM PLUMBI ACETATIS LEAD ACETATE OINTMENT

Lead Acetate, in fine powder, 20 grains, Paraffin Ointment, white, 480 grains (1 in 25)

Foreign Pharmacopœias—Official in Austri, Lead Acetate 1, Water 9, Wool Fat 45, Vaseline 45, Hung, Lead Acetate 3, Lard 100, White Wax 50, Water 10, Dan, Lead Acetate 1, Benzoated Lard 9, Norw, Lead Acetate 1, Olive Oil 14, Yellow Wax 5 Not in the others

Not Official

LOTIO PLUMBI ACETATIS—Lead Acetate, in powder, 2 grains, Diluted Acetic Acid, 2 minims, Distilled Water, up to 1 fl oz—*London Ophthalmic 1887*, omitted in 1901

This has been incorporated in the *B P C*

PLUMBI CARBONAS.

LEAD CARBONATE

$2\text{PbCO}_3, \text{Pb(OH)}_2$, eq 768.91

FR, CARBONATE DE PLOMB, GFR, CFRUSSA ITAL, CERUSSA;
SPAN, ALBAIALE DE O CTRUSA

An odourless and tasteless heavy white opaque powder, or in readily pulverisable masses

Solubility.—Insoluble in Water, soluble, with effervescence, in Diluted Nitric Acid and in Diluted Acetic Acid

Medicinal Properties.—Employed externally as an astringent and sedative, or as an ointment for ulcers and inflamed and excoriated surfaces

Official Preparation—*Unguentum Plumbi Carbonatis*

Foreign Pharmacopœias—Official in Austr, Hung and Jap (*Plumbum Carbonicum*), Ger and Swiss (*Cerussa*), Dan, Norw and Swed. (*Hydratocarbonas Plumbicus*), Dutch (*Carbonas Plumbicus*), Mex. (*Carbonato de Plomo*), Port (*Alvaiade*), Russ (*Plumbum Carbonicum Basicum*), Span (*Albayalde Cerusa*) Not in the others

Tests—Lead Carbonate is officially required to be completely soluble in diluted Acetic Acid, Carbon Oxide gas being simultaneously evolved. This gas when passed into Lime Water affords a precipitate soluble in a sufficient excess of the gas, and also redissolving in dilute mineral acid. The solution answers the tests distinctive of Lead given under *Plumbum*. It is official in the *P G*, but not in the *U S P*. The *P G* requires that it shall leave not less than 85 pc of Lead Oxide when ignited at a dull red heat. It may also contain Copper, Iron and matter insoluble in diluted Nitric Acid, such as siliceous material, Barium Sulphate, etc, in addition to Calcium, Magnesium or Zinc. Calcium, Magnesium and Zinc may be detected by applying the usual tests for these metals after the separation of the Lead from the acidified solution by Hydrogen Sulphide. Copper and Iron may be detected by the Potassium Ferrocyanide test described below. Siliceous matter may be detected by the residue remaining insoluble in a mixture of Nitric Acid and Water as described below.

Residue—The solution of the salt in Acetic Acid, after removal of the Lead by Hydrogen Sulphide gives a filtrate which, on evaporation to dryness, should not leave a weighable residue, *P G*.

If 1 gramme of the salt be dissolved in 2 cc of Nitric Acid with the addition of 4 cc of Water, not more than 0.01 gramme should be insoluble, *P G*.

Potassium Ferrocyanide—Sodium Hydroxide T.S. produces in the above solution a precipitate soluble in excess of the reagent. The same solution, with excess of Sulphuric Acid and filtered, gives a filtrate which should not be affected by T.S. of Potassium Ferrocyanide, *P G*.

Preparation

UNGUENTUM PLUMBI CARBONATIS. LEAD CARBONATE OINTMENT

Lead Carbonate, 1, Paraffin Ointment, white, 9

Foreign Pharmacopœias—Official in Austr, 3 in 10, Hung, Norw., Russ and Swed, 1 in 3, Dutch (*Unguentum Carbonatis Plumbici Camphoratum*) Camphor 3, Sesame Oil 5, Lead Carbonate 20, Lard 70, Mex (*Unguento Blanco*) simple and Port, 1 in 5, Ger (*Emplastrum Cerussæ*), Powdered Lead (simple) 5, Olive Oil 2, Lead Plaster 12

PLUMBI IODIDUM.

LEAD IODIDE

 PbI_2 , eq 457 15FR, IODURE DE PLOMB, GER, BLEIJODID, ITAL, JODURO DI PIOMBO,
SPAN, YODURO PLUMBICO

A golden yellow, odourless heavy crystalline powder Prepared by precipitating a soluble Lead salt with Potassium Iodide solution

The *USP* requires it to contain not less than 99 p.c of pure Lead Iodide It should be kept in well closed bottles of a dark amber tint and protected as far as possible from the light

Solubility—Sparingly soluble in cold Water, more soluble in boiling Water, soluble also in solutions of Acetates, and of Ammonium Chloride

Medicinal Properties—Used externally to reduce chronic inflammatory gland and joint enlargements, also in the form of pessaries

In 'dispersible' tumours of the mamma—*BMJ* '94, ii 972

Official Preparations—Emplastrum Plumbi Iodidi, and Unguentum Plumbi Iodidi

Not Official—Pessus Plumbi Iodidi, Pessus Plumbi Iodidi et Atropinae, and Pessus Plumbi Iodidi et Opii

Foreign Pharmacopœias—Official in Austr (Plumbum Iodatum), Fr (Iodure de Plomb), Ital (Ioduro di Piombo), Mex (Yoduro de Plombo), Port (Iodato de Chumbo), Swiss (Plumbum Iodatum); Span (Ioduro Plumbico), and US Not in the others

Tests—Lead Iodide dissolves in boiling Water, and the solution should answer the tests distinctive of Lead given under Plumbum. When strongly heated it evolves violet vapour The hot aqueous solution affords with Silver Nitrate Solution a curdy yellow precipitate insoluble in Nitric Acid, almost insoluble in Ammonia Solution, soluble in Potassium Cyanide Solution No method of determining the amount of pure Lead Iodide present in the sample is given in either the *BP* or *USP*

The more generally occurring impurities are Lead Chromate, insoluble foreign salts, Acetate, Nitrate and soluble matter of a foreign nature The *USP* includes a test for the absence of Acetate as described below with Hydrogen Sulphide and Ferric Chloride, and a test for the limit of Nitrate with Potassium Hydroxide Solution and Aluminium wire, which is also described in small type below Lead Chromate and insoluble foreign salts, if present, may be detected by the Ammonium Chloride test Soluble foreign salts, if present, may be detected by evaporating to dryness a portion of the filtrate after separation of the Lead by Hydrogen Sulphide, no residue should remain

Ammonium Chloride—On triturating 1 gramme of the salt with 2 grammes of Ammonium Chloride and 2 c.c of Water, a nearly white mixture will result, which, when heated in a test-tube on a water-bath for a few minutes, should give a clear and almost colourless solution, and if the above solution be

cooled a solid mass of nearly colourless fine silky crystals will be produced, and, on adding Water or diluted Sulphuric Acid to this mass, Yellow Lead Iodide will separate, *U S P*

Potassium Hydroxide Solution and Aluminium Wire—Add 0.1 gramme of the salt to 5 cc of Water, boil the mixture, cool and filter into a test-tube of about 40 cc capacity, then add 5 cc of Potassium Hydroxide T.S. and about 0.2 gramme of Aluminium wire, insert in the upper portion of the test-tube a pledget of purified Cotton and over the mouth place a piece of moistened red Litmus paper, then, if the tube be heated on a water bath for 15 minutes no blue coloration of the paper should be discernible, *U S P*

Hydrogen Sulphide Ammonia and Ferric Chloride—Boil 1 gramme of the salt with 20 cc of Water, cool and filter, remove the Lead from the filtrate by Hydrogen Sulphide and again filter. A portion of the second filtrate after boiling and carefully neutralising with Ammonia Water should not be coloured red by a drop of T.S. of Ferric Chloride, *U S P*

Preparations

EMPLASTRUM PLUMBI IODIDI. LEAD IODIDE PLASTER

Lead Iodide, 1, Lead Plaster, 8, Resin, 1 (1 in 10)

UNGUENTUM PLUMBI IODIDI. LEAD IODIDE OINTMENT

Lead Iodide, in fine powder, 1, Paraffin Ointment, yellow, 9 (1 in 10)

Foreign Pharmacopœias—Official in Fr, Mex, Port, Span and Swiss, 1 in 10. Not in the others.

An Ointment of Cadmium Iodide of the same strength has been recommended as a substitute, it is said not to stain the skin.

Not Official

PESSUS PLUMBI IODIDI—Lead Iodide, 5 grains, Oil of Theobroma, q s for one pessary

PESSUS PLUMBI IODIDI ET ATROPINÆ—Lead Iodide, 10 grains; Atropine Sulphate $\frac{1}{2}$ grain, (Gelatin) Basis, 60 grains

PESSUS PLUMBI IODIDI ET OPII—Lead Iodide, 5 grains, Opium, in powder, 2 grains. Oil of Theobroma, 12 grains

PLUMBI OXIDUM.

LEAD OXIDE

B P Syn—LITHARGE

PbO, eq 221.23

FR, OXYDE DE PLOMB FONDU, GFR, BLEIGLATT, ITAL, PROTOSSIDO DI PIOMBO, SPAN, LITARGIRIO

In odourless and tasteless, heavy yellow or reddish-yellow scales or powder, prepared by the atmospheric oxidation of molten metallic Lead

It should be kept in well-closed vessels. The *U S P* requires that it should contain not less than 96 pc of pure Lead Oxide. Neither the *B P* nor the *P G*. states a requisite percentage of pure Oxide.

Official Preparation.—*Emplastrum Plumbi* Used in the preparation of *Liquor Plumbi Subacetatis Fortis*, *Plumbi Acetas*, and *Glycerinum Plumbi*

Subacetatis Lead Plaster is contained in Emplastium Hydragryi, Emplastum Plumbi Iodidi, Emplastium Resinae, and Emplastium Saponis.

Not Official—Emplastium Lithargyri Compositum, Ung Diachylon Hebrae, Ung Diachylon Carbolisatum, Dr Pearson's Cerate, and Plumbi Oleas.

Foreign Pharmacopœias—Official in Austri, Hung, Russ and Swiss (Plumbum Oxydatum), Belg and Ger (Lythargyrium), Dan, Norw and Swed (Oxydum Plumbicum), Dutch (Oxydum Plumbicum Semivitreum), Fr (Oxyde de Plomb Fondu), Ital (Protossido di Piombo), Jap, Mex (Oxido de Plomo), Port (Oxydode Chumbo), Span (Litaigirio), U S (Plumbi Oxidum),

Tests—Lead Oxide dissolves in Acetic Acid or in Diluted Nitric Acid, and the resulting solution answers the tests distinctive of Lead given under Plumbum. When heated it becomes brownish-red fusing at a red heat, and when heated with reducing substances it leaves a residue of metallic Lead. Although a content of pure Oxide is specified in the *USP*, neither this Pharmacopœia nor the *BP* nor *PG* includes a method of determination.

The more generally occurring impurities are Copper, Iron and metallic Lead, Carbonates, Silicates, Barium Sulphate, impurities insoluble and soluble in Acetic Acid, excess of moisture. The presence of Copper and Iron may be detected by the Ammonia test given below. Metallic Lead, if present, is indicated by the formation of nitrous fumes when the sample is dissolved in Diluted Nitric Acid, the presence of Carbonates by effervescence during solution. Silica, Silicates, and Barium Sulphate, if present, remain insoluble.

In testing for impurities insoluble in Acetic Acid, both the *USP* and *PG* treat 5 grammes of the Oxide with 5 c c of Water, 20 c c of Acetic Acid (*USP*, 36 p c w/w, *PG*, 30 p c w/w), boiling the mixture for a few minutes and filtering from the insoluble residue, which, when well washed and dried, should amount to not more than 4 p c according to the *USP*, and not more than 0.1 p c according to the *PG*. In continuation of the *USP* test the mixed filtrate and washings from the insoluble matter are mixed with Hydrochloric Acid until a precipitate is no longer formed, the balance of the Lead is removed by Hydrogen Sulphide and the liquid filtered, the filtrate, when evaporated to dryness, should not leave more than 0.05 of a gramme, indicating not more than 0.1 p c of impurities soluble in Acetic Acid. When heated to a dull red heat in a porcelain crucible the sample, according to the *USP*, should lose not more than 4 p c, according to the *PG* 1 p c at the most, indicating the limit of moisture and Carbonate allowed by the respective Pharmacopœias.

Ammonia—The solution of Lead Oxide in Nitric Acid after the addition of excess of Sulphuric Acid and filtration gives a filtrate which, when supersaturated with Ammonia Water, should only assume a slight bluish tint and yield only traces of a reddish-brown precipitate, *PG* and *USP*.

Preparation

EMPLASTRUM PLUMBI. LEAD PLASTER *NO Syn*—DIACHYLON PLASTER

Lead Oleate with mechanically included Glycerin, obtained by boiling together Lead Oxide, Olive Oil, and Distilled Water

Equal weights of Lead Plaster and Soap Plaster melted together form an excellent plaster for coins

Foreign Pharmacopœias—Official in Austr, Litharge 1, Sesame Oil 1, Lead 1, Belg, Litharge 2, Olive Oil 2, Water 1, Lead 2, Dan, Litharge 5, Olive Oil 10, Water 1, Dutch, Pot and Russ, Litharge 1, Lead 1, Olive Oil 1, Water *q s*, Fr, Litharge 1, Lead 1, Olive Oil 1, Water 2, Ger and Jap, Litharge 5, Olive Oil 5, Lead 5, Water 1, Hung, Litharge 1, Lead 2, Ital, Litharge 1, Water 1, Olive Oil 2, Mex (Emplasto Simple), Litharge 2, Lead 4, Water 3, Nor and Swed, Litharge 1, Olive Oil 2, Water *q s*, Span, Litharge 1, Olive Oil 2, Water 2, Swiss, Litharge 16, Olive Oil 30, Water *q s*, U S, Lead Acetate 60, Soap 100, Water *q s*

Not Official

EMPLASTRUM LITHARGYRI COMPOSITUM (Ger.)—Lead Plaster 12, Yellow Wax 1½, Gum Ammoniacum 1, Galbanum 1, Turpentine 1

Foreign Pharmacopœias—Official in Belg (Emplastum Diachylon Gummosum), Lead Plaster 72, Yellow Wax 7, Gum Ammoniacum 7, Galbanum 7, Turpentine 7, Fr (Emplâtre Bism) Olive Oil 10, Lead 5, Butter 5, Yellow Wax 5, Litharge 5, Purified Mutton Suet 5, Purified Black Pitch 1, also Emplâtre Diachylon Gomme, Litharge 62, Lead 62, Olive Oil 62, Water 125, Yellow Wax 12, Purified Burgundy Pitch 12, Venice Turpentine 12, Purified Ammoniacum 10, Purified Galbanum 10, Oil of Turpentine 6, Ger (Emplastrum Fuscum Cimphorum), Red Lead or Vermilion 30, Olive Oil Commune 60, Yellow Wax 15, Camp. 1, U S, Ital (Emplastro Diachylon Gommosinoso), Lead 1, Yellow Wax 1, Gum Ammoniacum 1, Galbanum 1, Turpentine 1, Jap, Lead Plaster 12, Yellow Wax 13, Gum Ammoniacum 1, Galbanum 1, Turpentine 1, Span (Emplasto de Plomo Gomado), Turpentine 60, Yellow Wax 85, Galbanum 85, Gum Ammoniacum 85, Lead Plaster 735, Swiss (Emplastum Plumbi Compositum), Lead Plaster 72, Yellow Wax 9, Ammoniacum 6, Galbanum 6, Turpentine 7, Water *q s*

UNG DIACHYLON HEBRÆ (modified by Professor Kaposi)—Simple Lead Plaster 1, Soft Paraffin 1, melt with heat

Unguentum Diachylon (*Hebra*) according to Hager, Hebra's original formula for this ointment was equal parts of Lead Plaster and Linseed Oil, and this formula is in the Gross Hospital Pharm (1884), but the majority of the Hospitals, including Charing Cross (1904), are made with Soft Paraffin or a mixture of Hard and Soft Paraffin whilst some employ Olive Oil. It is also known as **Unguentum Plumbi Oleatis**

Foreign Pharmacopœias—Official in Austr (Unguentum Plumbi Oxydati), Litharge 20, Sesame Oil 40, Lead 40, to make Unguentum Diachylon Hebra add 2 pc of lavender Oil, Dan (Unguentum Oxydi Plumbici), and Swed (Unguentum Diachylon), Lead Plaster 13, Liquid Paraffin 7, Dutch (Unguentum Diachylon), Lead Plaster 1, Sesame Oil 1, Ger and Russ (Unguentum Diachylon), Lead Plaster 1, Olive Oil 1, Jap (Unguentum Hebra), Lead Plaster 1, Olive Oil 1, Mex (Unguentum Inducativo), Red Oxide of Lead 6, Lead 50, Swiss (Unguentum Hebræ), Lead Plaster 50, White Vaseline 43, Glycerin 7, Ethereal Tincture of Benzoin 5, U S, Lead Plaster 50, Olive Oil 49, Lavender Oil 1

UNGUENTUM DIACHYLON CARBOLISATUM—Liquid Carbolic Acid 1, Diachylon Ointment 49

DR. PEARSON'S CERATE—Lead Plaster 4, Yellow Beeswax 1, Oil of Almonds 3. Melt and mix

PLUMBI OLEAS—Lead Acetate, 250 grains, dissolve in Distilled Water, 40 fl oz, add slowly Solution of Sodium Oleate (1 Castile Soap in 20), 20 fl oz., warm gently, wash by decantation, collect and dry

Melted with equal parts of Lard, and Lard Oil or Olive Oil, to form an ointment

PLUMBI SUBACETATIS LIQUOR FORTIS.

STRONG SOLUTION OF LEAD SUBACETATE

B P Syn—GOULARD'S EXTRACT

FR., ACETATE BASIQUE DE PLOMB DISSOUS, GFR., BLEI-SSIG, ITAL., ACETATO BASICO DI PIOMBO, SPAN., ACETATO PLUMBICO LIQUIDO

A clear colourless heavy liquid, having a sweet astringent taste. It is a solution of Lead Subacetate, $Pb_2O(C_2H_3O_2)_2$, eq 543.74 in Water, and is prepared by boiling 5 of Lead Acetate, and $3\frac{1}{2}$ of powdered Lead Oxide in 20 of distilled Water for 30 minutes, maintaining the volume of the liquid by addition of Distilled Water, filtering and making up to 20 with Distilled Water.

Medicinal Properties—When largely diluted, as in Lotio Plumbi Evaporans or as Lotio Plumbi Evaporans cum Morphina, it is used externally as an astringent and sedative for inflammation arising from sprains, bruises, etc. Sometimes used as an astringent gargle ($\frac{1}{2}$ fl drim to 6 fl oz of Rose Water). A good astringent application to external piles is—Strong Solution of Lead Subacetate, 2 to 3 fl drim, Solution of Morphine Acetate, 3 fl drim, Distilled Water, to 6 fl oz.

Incompatibles—Hard Water, mineral Acids, vegetable Acids, Alkalis, Chlorides, Iodides, all astringents, preparations of Opium, and Mucilage of Acacia.

Official Preparations—Glycerinum Plumbi Subacetatis, Liquor Plumbi Subacetatis Dilutus, and Unguentum Glycerini Plumbi Subacetatis.

Not Official—Ceratum Plumbi Compositum, Cremor Lithargyri, Lotio Plumbi, Lotio Plumbi cum Opio, Lotio Plumbi Evaporans, Lotio Plumbi Evaporans cum Morphina, Lotio Plumbi Lactatis, and Unguentum Plumbi Tannici.

Antidotes—Wash out the stomach or give an emetic, Sodium or Magnesium Sulphate, liberal libations of Milk, or White of Egg mixed with Water, Opium or Belladonna in Lead colic.

A course of Potassium Iodide is useful in eliminating Lead from the system.

L '81, ii 779 gives an unusual source of Lead poisoning, viz, shot found in a bottle full of Port Wine, an appreciable quantity of Lead was found in solution.

Foreign Pharmacopœias—Official in all, U.S. sp gr about 1.235 at 25° C (77° F) (Plumbum Aceticum Basicum Solutum)—Aust. and Hung. sp gr 1.230 to 1.240, Russ. sp gr 1.235 to 1.242, Belg. (Subacetatis Plumbi Liquidus), sp gr 1.240, Dan. Norw. and Swed. (Solutio Subacetatis Plumbici), sp gr 1.165 to 1.170, Dutch (Solutio Acetatis Plumbici Basici), sp gr 1.235 to 1.240, Fr. (Acetate Basique de Plomb Dissous), sp gr 1.320, Ger. (Liquor Plumbi Subacetici), sp gr 1.235 to 1.240, Ital. (Acetato Basico di Piombo), sp gr 1.32, Jap. sp gr 1.23 to 1.24, Mex. (Acetato de Plomo Liquido), sp gr not given, Port. (Solutio de Subacetato de Chumbo), sp gr 1.260, Spain. (Acetato (sub) Plumbico Liquido), sp gr 1.32, Swiss (Plumbum Subaceticum Solutum), sp gr 1.235 to 1.240.

Tests—Strong Lead Subacetate Solution when freshly prepared has a sp gr of 1.277. It is officially required to possess a gravity of 1.275. The Liquor official in the *P G* has a sp gr from 1.235 to 1.240, that of the *U S P* about 1.235 at 25° C (77° F). It possesses an alkaline reaction towards red Litmus paper, the *P G* states that it does not redden Phenolphthalein Solution. The *B P* states that it

forms an opaque white jelly with Gum Acacia Mucilage, the *U S P* that it affords a dense white precipitate when added to Acacia Solution. It answers the tests distinctive of Lead given under Plumbum. When treated with Sulphuric Acid it evolves a strong acetous odour, and when warmed with Acetic Acid and Alcohol (90 p c) a characteristic odour of Ethyl Acetate is evolved. It is officially required to contain 23.1 p c w/w of pure Lead Subacetate as determined by the process indicated below. The *U S P* requires that it shall contain in not less than 25 p c w/w of Lead Subacetate as volumetrically determined by the process indicated below. The Liquor should be free from the impurities given under Lead Acetate. The Subacetate may be distinguished from the Normal Acetate by the test with Acacia Mucilage above.

The *P G* includes a test for Iron as given below with Potassium Ferrocyanide Solution. As the Liquor has a tendency to absorb Carbon Dioxide from the air, it should be kept in well-closed glass bottles and exposed as little as possible.

Potassium Ferrocyanide—After the addition of Acetic Acid, solution of Lead Subacetate should give, with Potassium Ferrocyanide T S, a pure white precipitate, *P G*.

Volumetric Determination—Dilute 10 grammes of the solution to 100 c c with previously boiled and cooled Distilled Water, take 13.6 (13.595) c c of this and add it to 35 c c of Tenth-normal Volumetric Oxalic Acid Solution in a graduated cylinder, shake thoroughly and dilute the mixture to 50 c c with Distilled Water, and again shake. After the precipitate has settled 10 c c of the clear solution diluted with about 50 c c of Water and 5 c c of Tenth-normal Potassium Permanganate Solution to produce a permanent pink tint, *U S P* 17 c c of Tenth-normal Volumetric Sulphuric Acid Solution should be required to completely precipitate 1 gramme of the Strong Lead Subacetate Solution, *B P*.

Preparations

GLYCERINUM PLUMBI SUBACETATIS. GLYCERIN OF LEAD SUBACETATE

Lead Acetate, 5, Lead Oxide, in powder, $3\frac{1}{2}$, Glycerin, 20, Distilled Water, 12 Min. Boil for a quarter of an hour, filter, evaporate at a temperature not exceeding 222° F (105.5° C) until the product weighs $32\frac{1}{2}$, and has a sp gr of 1.48.

This is more conveniently made with half the quantity of the Distilled Water.

Foreign Pharmacopœias—Official in Port, Solution 1, Glycerin 9. Not in the others.

Glycerin Lead Subacetate has a sp gr of 1.480 to 1.485. The *B.P.* figure 1.48.

LIQUOR PLUMBI SUBACETATIS DILUTUS. DILUTED SOLUTION OF LEAD SUBACETATE *B P Syn*—GOULARD'S LOTION; GOULARD WATER. *N O Syn*—AQUA VEGETO-MINERALIS GOULARDI, AQUA SATURNINA, AQUA DE VEGETO.

Strong Solution of Lead Subacetate, 2 fl drms, Alcohol (90 p.c.), 2 fl drms, Distilled Water, q s. to make 20 fl oz. (1 in 80)

As the diluted Liquor is liable to absorb Carbon Dioxide from the air, it should be kept in well-closed bottles and exposed as little as

possible The *BP* solution is not specifically required to contain any definite percentage of Lead Subacetate, the *USP* must contain about 1 p c The *PG* does not include a dilute Liquor

Foreign Pharmacopœias—Official in Austr (Aqua Goulardi), Solution 2, Spirit of Wine (60 p c) 5, Water 93, also (Aqua Plumbica), Solution 1, Water 49, Dan and Norw (Aqua Saturnina), Dutch (Aqua Plumbi Goulardi) and Swed (Solutio Subacetatis Plumbici Diluta), Solution 2, Diluted Alcohol 8, Water 90, Fr (Lotion à l'Acetate de Plomb), Solution 1, Water 50, Ger and Swiss (Aqua Plumbi), Solution 1, Water 49, Ital (Aqua Saturnina), Solution 1, Water 50, Hung (Aqua Goulardi), Solution 2, Alcohol (70 p c) 5, Water 100, also (Aqua Plumbica), Solution 1, Water 50, Jap (Liquor Plumbi Subacetici Dilutus), Solution 2, Water 98, Mex (Agua de Vegeto), Solution 3, Eau de Cologne 5, Water 92, Port (Aqua Saturnina Alcoholisata), Solution 2, Alcohol (85 p c) 8, Water 90, also (Aqua Saturnina), Solution 1, Water 50, Russ (Aqua Plumbi Spirituosa), Solution 2, Water 98, also (Aqua Plumbi), Solution 1, Water 49, Span (Agua Vegeto Mineral), Solution 1, Alcohol (95 p c) 2, Distilled Water 97, U S, Solution 4, Water to make 100

Tests.—Diluted Lead Subacetate has a sp gr of 1.002 No official method is given for determining the amount of Lead Subacetate present

UNGUENTUM GLYCERINI PLUMBI SUBACETATIS. LEAD SUBACETATE OINTMENT

Glycerin of Lead Subacetate (by weight), 1, Paraffin Ointment, white, 5

Foreign Pharmacopœias—Official in Belg (Unguent Subacetatis Plumbi), 8 in 10, Dutch (Ung Plumbici Basici), 1 in 4, Ital (Pomata con Acetato di Piombe), Ger and Swiss (Unguentum Plumbi), 1 in 10, Russ (Ung Plumbi Acetici), 1 in 12, Swed (Ung Subacetatis Plumbici), 8 in 20, U S (Ceratum Plumbi Subacetatis), 1 in 5 Not in the others

Not Official

CERATUM PLUMBI COMPOSITUM—Solution of Diacetate of Lead, 6 fl oz, Beeswax, 8 oz, Olive Oil, 20 fl oz, Camphor, 1 dr m —*PL*

This has been incorporated in the *BPC* as follows —

Camphor, 0.5, Yellow Beeswax, 23.5, Olive Oil, 58.5, Solution of Lead Subacetate, 17.5

CREMOR LITHARGYRI—Solution of Lead Subacetate, 1, Cream, 7 Mix Useful in eczema

LOTIO PLUMBI—Strong Solution of Lead Subacetate, 2 fl dr m, Water, q s to make 20 fl oz —*St Thomas's*

This has been incorporated in the *BPC*

LOTIO PLUMBI CUM OPIO—Tincture of Opium, 20 minims; Lead Lotion, to 1 oz —*Lock*

Tincture of Opium, 5, Lead Lotion, to 100 —*BPC*

LOTIO PLUMBI EVAPORANS—Strong Solution of Lead Acetate, 2 fl dr m, Rectified Spirit, 1½ fl oz, Rose Water, to 8 oz —*Squire*

Strong Solution of Lead Subacetate, 2 fl dr m, Alcohol (90 p c), 4 fl oz, Water, q s to make 20 fl oz —*St Thomas's*

This has been incorporated in the *BPC*

LOTIO PLUMBI EVAPORANS CUM MORPHINA—The Solution given above, 7 fl oz, Solution of Morphine Acetate, 1 fl oz —*Squire*

This is an improvement on the old Lead and Opium Solution, with its coloured deposit of Lead Meconate

LOTIO PLUMBI LACTATIS—Solution of Lead Subacetate, 1 fl drm ; Salicylic Acid, 2 grains, Milk, to 2 fl oz—*Middlesex*

UNGUENTUM PLUMBI TANNICI—*Ger*, Tannic Acid 1, Liquor Plumbi 2, Lard 17

Hung and *Swiss*, Tannic Acid 1, Liquor Plumbi 2, Vaseline 17

Prus, Tannic Acid 1, Glycerin 3, Liquor Plumbi 6, Ung Cerat 24

Sind, Tannic Acid 1, Subacetate of Lead Solution 3, Wool Fat 3, Vaseline 3

PODOPHYLLI RHIZOMA.

PODOPHYLLUM RHIZOME

B P Syn—PODOPHYLLUM ROOT

FR PODOPHYLLI GER, PODOPHYLLIN, ITAL, PODOFILLO, SPAN, PODOFILO.

The dried Rhizome and Roots of *Podophyllum peltatum*, L. Imported from North America

The dried Rhizome and Roots of *Podophyllum Emodi*, L. and the Resin extracted from the same dose $\frac{1}{2}$ to 1 grain = 0.016 to 0.06 gramme, are official in the *Ind* and *Col* but for India and the Eastern Colonies

The Resin obtained from *P Emodi* is as valuable a purgative as that obtained from *P peltatum*

Medicinal Properties.—The Resin is an active cholagogue and, in large doses, purgative, in doses of $\frac{1}{8}$ to $\frac{1}{4}$ grain it is a common ingredient of pills for habitual constipation associated with liver disorder

Prescribing Notes—The Resin is given in pills, combined with Extract of Henbane or Belladonna to prevent griping, a few grains with a Aloes or Colocynth, sometimes $\frac{1}{2}$ grain of each. In its nature, the addition of Aromatic Spirit of Clove in Tinctura Podophylli Ammonata, since Water does not precipitate the Resin from this and it does so from the ordinary Tincture

Official Preparations—Podophylli Resina and Tinctura Podophylli

Not Official—Ort R.

Podophylli Compositæ Pil.

Aloes et Podophylli Compositæ, Pilule Podophylli Belladonnae et Capsici, Pilule de Podophylline Belladonnae Tinctura Podophylli Ammonata

Foreign Pharmacopœias—Official in Belg, Dutch, Fr, Ital, Port, Spain and U S. Not in the others

Descriptive Notes.—Podophyllum Rhizome as met with in commerce occurs in cylindrical pieces, 1 to 6 or 8 in in length, and 2 to 4 lines in diameter (5 to 8 mm, *B P.*), of a chocolate or reddish-brown colour, marked on the upper surface, at intervals of 2 or 3 in, with the circular scars of former stems, and on the under surface near the nodes, with little rootlets about half a line thick, which are frequently more or less broken off. If shrunken, wrinkled, and flattened, the Rhizomes are of inferior quality. The fracture is short, normally mealy, but hoary if overheated in drying; it exhibits a thin bark, a pith 2 lines in diameter in the larger pieces, and a thin circle of 20–40 vascular bundles. It has an acrid, bitter taste and a characteristic odour.

The Rhizome of *Podophyllum Emodi* has been used as a source of

Podophylli Resina It is quite different in appearance from that of *Podophyllum peltatum*, consisting of a compressed knotty Rhizome about $\frac{1}{2}$ in broad (12 mm), and $\frac{1}{4}$ in (8 mm) thick, the upper surface covered with contiguous circular stem scars, the whole under surface having brown or blackish simple roots about $\frac{1}{12}$ to $\frac{1}{8}$ in (2 to 3 mm) thick, with short branches

Tests—*Podophyllum* Root contains from 3 to 4 p c of ash, and should not contain more than 5 p c. A standard of not less than 50 p c of Resin soluble in Alcohol (90 p c) has been suggested (*YBP* '03, 247, *PJ* '02, 11 496). American *Podophyllum* Rhizome may contain from 4 to 6 p c of *Podophyllum* Resin, the Indian Rhizome from 10 to 12 p c. Seven samples of the root examined (*PJ* '03, 1 164) yielded percentages of Resin varying from 1.6 to 3.86, an average of 2.19 p c.

Preparations

PODOPHYLLI RESINA *PODOPHYLLUM RESIN* *NO Syn*—**PODOPHYLLIN**

The Resin is extracted by Alcohol (60 p c), the solution concentrated in a still, and the residue poured into acidulated Water to precipitate the Resin, which is washed and finally dried at a temperature not exceeding 100° F (37.7° C).

A yellow, greenish-yellow, or yellowish-brown, amorphous powder, or amorphous masses readily reduced to powder. It has a faint, peculiar odour, and a bitter taste.

The variations in colour appear to depend upon the heat applied during its preparation, by distilling quickly and drying at a low temperature the lightest tints are obtained.

It should be kept in well-closed glass receptacles of a dark amber tint in a cool atmosphere and protected as far as possible from contact with the air and light.

The *BP* Resin is obtained from the dried Rhizome and Roots of *Podophyllum peltatum*, the Resin obtained from the dried Rhizome and Roots of *Podophyllum Emodi* is official in the *Ind* and *Col Add*. The *USP* and the *PG* only admit the Resin obtained from *Podophyllum peltatum*.

Dose.— $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme

Not Official—See *Podophylli Rhizoma*

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Ger, Hung, Russ and Swiss (*Podophyllinum*), Dutch, Jap, Norw, Port, Swed and US (*Resina Podophylli*), US has also a Fluid Extract, Fr (*Resine de Podophylle*), Ital (*Podofillina*), Mex and Span (*Podofilina*).

Tests.—*Podophyllum* Resin darkens in colour when heated or when exposed to the light. It is required by the *USP* to be soluble or nearly so in Alcohol (90 p c) and in Ammonia Solution, to be reprecipitated from its solution in Alcohol (90 p c) by Water, and from its solution in Ammonia Solution on acidification. It is also

officially required to be partly soluble in Ether. It is difficult to find a commercial sample perfectly soluble in cold Alcohol (90 p c), and many will not give clear solutions even with the addition of Ammonia. The amount of matter insoluble in Alcohol (90 p c) should, however, not exceed 5 p c. The *USP* states that not less than 99 p c. of Resin should be soluble in Alcohol (94.9 p c), and this statement appears to be based upon the results of some experiments by Goudin and Merrell recorded (*Proc Amer Pharm Assoc* '02, 348). Gravell and Sage [*PJ* (3) xxiv 421] also state that a good sample should dissolve almost completely in Rectified Spirit, *BP* 1885. The *PG* states that it is soluble in 10 parts by weight of Alcohol (90 p c) to a brown fluid which is precipitated by the addition of Water. A good sample of the *Peltatum* Resin should dissolve almost completely in Ammonia, and not more than 5 p c should remain insoluble.

The *Emodi* Resin is stated to gelatinise with Ammonia Solution, the gelatinisation being stated (*CD* '03, 1 630) to be due to the fact that *Podophyllum Emodi* contains from 1½ to 3 times as much Podophyllotoxin as the *Podophyllum peltatum*.

Dott (*PJ* '06, ii 431, *Proc Amer Pharm Assoc* '07, 681) gives an Ammonia test for Podophyllin, and he applies the test by treating 0.5 of a gramme of the Resin with 30 cc of equal volumes of Liquor Ammoniae and Water, stirring and bringing well into contact for 5 minutes, filtering the liquor through a counterpoised filter, washing with Water until the washings are practically colourless, drying till constant, then weighing. He finds the *Emodi* Resin remains practically insoluble, whilst the residue from *Peltatum* should not amount to more than 15 p c of its original weight. The Report of the Committee of Reference in Pharmacy states that the Committee does not know a satisfactory test to distinguish the *Podophyllum Emodi* from *Podophyllum peltatum*. The solubility in Ammonia Solution is not considered (*PJ* '02, ii 368) of much value. The *USP* does not include a statement as to the solubility in Ammonia Solution. The *PG* states that *Podophyllum* should dissolve in 100 parts of Ammonia Solution to a yellowish-brown fluid which is miscible with Water, but which is again precipitated as a brown flocculent precipitate on the neutralisation of the Ammonia. The *USP* states that not less than 75 p c should be soluble in Ether, the *PG* that it is only partly soluble in Ether. The *USP* limit of 75 p c soluble in Ether seems very high, the usual amount averaging about 60 p c. The *BP* makes no reference to the solubility in Chloroform. The *USP* requires that not less than 65 p c should be soluble in Chloroform. The *PG* does not refer to the Chloroform solubility. More than half the weight of the Resin should dissolve in cold Chloroform, the residue being generally reckoned as medicinally inert, if the Chloroform solution be evaporated to a small bulk and poured into an excess of Ether, another inert body (Podophyllic Acid) is precipitated. If the Ether-chloroform Solution be now added to a large excess of Petroleum Ether there is precipitated a compound called Podophyllotoxin, supposed to contain the whole medicinal elements of the Resin.

From the results recorded (*P J* '02, 11 368) of the examination of a number of specimens of the Resin it is concluded that the limit of 50 p c soluble in Chloroform given by Squire is a good criterion of the quality of the sample. It is also suggested in the same reference that at least 40 p c of the original Resin should be precipitated from the chloroformic solution by Petroleum Ether. A method of determining the crude Picro-podophyllin has been suggested (*Proc Amer Pharm Assoc* '02, 346), it consists in treating 5 grammes of Podophyllin Resin, in a strong round bottle holding about 200 c c, with 10 grammes of freshly prepared Calcium Hydride, the bottle is closed with a good cork and the whole weighed, it is uncorked, transferred to a water-bath heated to 60° to 65° C (140° to 149° F) for a few minutes and 15 c c of Alcohol poured in, the bottle stoppered, well shaken, replaced in the water bath and retained there stoppered for 8 hours, shaking at first every few minutes to prevent the formation of a hard lump, after half an hour it is only necessary to shake the mixture about every quarter of an hour. The bottle is then cooled, about 7 c c of Chloroform added, it is placed on the balance and sufficient of a mixture of 2 parts by volume of Alcohol (94.9 p c) and 1 part by volume of Chloroform poured into the bottle to make the whole liquid added weigh 130 grammes. The bottle is shaken for a few minutes, set aside until the supernatant liquid becomes perfectly clear, and 65 grammes of the clear liquid are drawn off into a tared vessel, evaporated to dryness, the residue dried till constant in weight, and weighed. The percentage of crude Picro-podophyllin found varied from 15 to 22 p c, averaging 20 p c.

The *USP* states that not more than 25 p c should be dissolved in boiling Water. Neither the *BP* nor the *PG* makes any reference to the Water solubility. The *USP* also states that the hot aqueous solution deposits most of its content on cooling, and after the cooled liquid be filtered, the filtrate has a bitter taste and yields on the addition of a few drops of Ferric Chloride T S a brown coloration. It is soluble in Potassium or Sodium Hydroxide T S with the production of a deep yellow liquid which gradually becomes darker on standing. On neutralisation of the Potassium or Sodium Hydroxide the Resin is reprecipitated. The behaviour with Liquor Potassæ is stated to form a useful test for differentiating the *Peltatum* and *Emodi* Resin. 6 grains of the Resin should be mixed with 1 fl drim of diluted Alcohol and 8 or 10 drops of Liquor Potassæ. The *Peltatum* Resin should form a clear deep yellow liquid on shaking, the *Emodi* Resin becomes a semi-solid gelatinous mass. The alcoholic solution of the Resin should be only faintly acid in reaction towards blue Litmus paper. It is officially required to yield not more than 1 p c of ash when ignited with free access of air. Badly adulterated specimens are frequently detected by high percentage of ash, it may be as low as $\frac{1}{2}$ p c and should not exceed 1 p c. The *USP* states that it should not yield more than 1 p c of ash. The *PG* does not give an ash limit. The *BP* limit of 1 p c is generally considered a suitable one.

TINCTURA PODOPHYLLI.—TINCTURE OF PODOPHYLLUM

Podophyllum Resin, 320 grains, Alcohol (90 p c), *q. s.* to yield 20 fl oz of filtered product

Dose.—5 to 15 minims = 0.3 to 0.9 c c

15 minims equals $\frac{1}{2}$ grain of Podophyllum Resin. It is twice the strength of *B P '85*

A corresponding preparation, *Tinctura Podophylli Indiciæ* (1 in 30), dose 5 to 15 minims = 0.3 to 0.9 c c, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Tests.—Tincture of Podophyllum has a sp gr of 0.840 to 0.850, it contains about 3.5 p c of w/v total solids and about 87 p c w/v of Absolute Alcohol. A standard of 3.6 p c has been suggested

Not Official

PILULA PODOPHYLLI COMPOSITA—Podophyllum Resin, $\frac{1}{2}$ grain, Quinine Sulphate, 1 grain, Alcoholic Extract of Belladonna, $\frac{1}{2}$ grain, Extract of Socotrine Aloes, 1 grain — *University*

This has been incorporated in the *B P C* under the title *Poore's pill*

Podophyllum Resin, $\frac{1}{2}$ grain, Mercurous Chloride, 1 grain, Alcoholic Extract of Belladonna, $\frac{1}{2}$ grain — *St Thomas's*

This has been incorporated in the *B P C*

PILULÆ ALOES ET PODOPHYLLI COMPOSITÆ *Syn* Janoway's Pills—Purified Aloes, 1 grain, Resin of Podophyllum, $\frac{1}{2}$ grain, Extract of Belladonna Leaves, $\frac{1}{2}$ grain, Extract of Nux Vomica, $\frac{1}{2}$ grain in each pill — *USNF*

Pilulæ Podophyllini et Belladonnæ et Nuc Vomiciæ *Pharmacopœia* Resin, $\frac{1}{2}$ grain, Alcoholic Extract of Belladonna Leaves, $\frac{1}{2}$ grain, Extract of Barbadoes Aloes, 1 grain in each pill — *B P C*

Pilulæ Aloini et Podophylli Compositæ—Aloin, 4 grains, Capsicum 2 grains, Jalap Resin, 4 grains, Podophyllum Resin, $\frac{1}{2}$ grain, Extract of Nux Vomica, 2 grains, Green Extract of Hyoscyamus, 2 grains in each 10 pills — *B P C*

PILULÆ PODOPHYLLI, BELLADONNÆ ET CAPSICI—Resin of Podophyllum, 1.6, Extract of Belladonna Leaves, 0.8, Capsicum, 3.2, Sugar of Milk, 6.5, Acacia, 1.6, Glycerin, Syrup, each a sufficient quantity to make 100 — *USP*

PILULES DE PODOPHYLLINE BELLADONÉES *Pharmacopœia* 3 grammes, Extract of Belladonna, 0.1 gramme, Medicinal Syrup, $\frac{1}{2}$ ounce, make into 10 pilules — *F*

TINCTURA PODOPHYLLI AMMONIATA *Pharmacopœia* 24 grains, Alcohol (90 p c), 2 fl oz, Solution of Ammonia, 1 fl oz

Tests—Ammoniated Tincture of Podophyllum has a sp gr of 0.906, contains about 1.5 p c w/v of total solids and about 56 p c w/v of Absolute Alcohol

As the Resin does not separate on the addition of Water, this tincture is miscible with Water

Dose—10 to 30 minims = 0.6 to 1.8 c c. 1 fl dm contains 1 grain of the Resin

Podophyllum, 1; Aromatic Spirit of Ammonia, 50, dissolve, and after standing decant.—*Martindale*

This has been incorporated in the *B P C*

Not Official POTASSIUM

POTASSIUM

K, eq 38.83

Potassium was discovered by Sir Humphry Davy in 1807. It is a soft metal, cutting like wax, of a silver white colour, but tarnishes the instant it is cut, and assumes a leaden colour.

Tests—Potassium has a sp gr of 0.865. When freshly cut has a silver white lustre, but rapidly absorbs oxygen from the air and assumes a leaden colour. When a pellet is thrown upon water, hydrogen is set free, the heat developed during the action being so great that the evolved hydrogen is ignited. The resulting solution possesses a strongly alkaline reaction towards red litmus paper. A trace of a potassium salt, when moistened with hydrochloric acid and inserted into a non-luminous flame, imparts to it a distinctive violet coloration, which has a reddish violet tinge when viewed through blue glass, the yellow colour imparted to the flame by sodium compound is obscured by blue glass. A solution of a potassium salt, preferably a chloride or one in which hydrochloric acid is present, affords if sufficiently concentrated a yellow crystalline precipitate with platinum chloride solution, yielding upon ignition a residue of metallic platinum and potassium chloride. If the residue be dissolved in a little water, acidified slightly with nitric acid the solution yields on the addition of silver nitrate a white curdy precipitate, insoluble in nitric acid, soluble in ammonia solution. The aqueous solution, if sufficiently concentrated, yields on the addition of tartaric acid a white crystalline precipitate of potassium hydrogen tartrate; acetic acid or sodium acetate is added when the potassium is combined with a mineral acid. The best general reagent for potassium salts is probably a saturated solution of picric acid, a 1 p.c. solution of potassium nitrate yields a crystalline precipitate after a few seconds' shaking, whereas with tartaric acid no reaction is obtainable in 4 hours. Potassium salts may be distinguished from ammonium salts by the behaviour with platinum chloride solution, the precipitate from a solution of a potassium salt yielding on ignition, as above stated, a residue of potassium chloride and metallic platinum, a precipitate from an ammonium salt yielding on ignition a residue of metallic platinum only. Potassium salts may be distinguished from sodium salts by the latter not yielding a precipitate with platinum chloride solution, and by the violet colour imparted to the flame by the former, whereas the latter impart a strong yellow colour, also solutions of sodium salts yield no crystalline precipitate with tartaric acid.

The prolonged use of potassium salts is apt to lead to a depressant effect on muscular tissue, including that of the heart, in people with weakness of that organ this should be borne in mind.

POTASSA CAUSTICA.

POTASSIUM HYDROXIDE

B.P. Syn—CAUSTIC POTASH, POTASSIUM HYDRATE
HYDRATE OF POTASSIUM, *B.P.* '85

FR, HYDROXYDE DE POTASSIUM OFFICINAL, GFR, KALIUMHYDROXID,
ITAL, POTASSA CAUSTICA, SPAN, HIDRATO POTASICO

White, deliquescent sticks or pencils, or in hard, white, or nearly white, deliquescent cakes, officially stated to consist of potassium hydroxide, KOH, eq 55.71, with not more than 10 p.c. of combined water and impurities.

On account of its intense causticity and strong action on organic tissues great caution should be used in handling it. As it rapidly absorbs both carbon

Dioxide and moisture, and should be preserved from the air in well-closed hard glass bottles.

Commercial Caustic Potash as a rule contains 1 or 2 p.c. of Chloride derived from the Carbonate used in its preparation. When required *pure* it is dissolved in Absolute Alcohol, and the solution evaporated as far as practicable without access of air to avoid absorption of Carbonic Acid. No commercial samples, however, are quite free from Carbonate.

Solubility.—2 in 1 of Water, 1 in $3\frac{1}{2}$ of Alcohol (90 p.c.), 1 in 3 of Glycerin, 1 in 4 of Alcohol (60 p.c.) (if stronger than this the Alcohol separates).

Medicinal Properties.—A powerful caustic. Has been used for the destruction of tumours and to stimulate ulcers.

Prescribing Notes.—*It has a great tendency to burn, and attach the surrounding tissues, its action should be carefully guarded by means of Vaseline or Sticking Plaster. When mixed with Lime, as in 'Vienna Paste' (see p. 931), it is more easily controlled.*

Official Preparation.—Liquor Potassæ, used in the preparation of Potassii Permanganas.

Not Official.—Brandish's Alkaline Solution, and Potassa cum Calce (Vienna Paste).

Foreign Pharmacopœias.—Official in Austr and Hung (Kalium Hydro-oxidum), Austr (Kalium Hydro-oxidum Caustica Fusa), Dan, Dutch, Norw and Swed (Hydras Kalicus), Fr (Hydroxyde de Potassium Officinal, also Hydroxyde de Potassium Ordinaire), Ger and Russ (Kali Causticum Fusum), Ital, (Potassa Caustica), Jap (Kali Causticum), Mex (Oxido de Potasio), Port (Hydrato de Potassa), Span (Hidrato Potasico), also (Potassa Caustica por la Cal); Swiss (Kalium Hydroxidum), US (Potassii Hydroxidum).

Tests.—Potassium Hydroxide fuses when strongly heated, the *U.S.P.* states when heated to a temperature of about 530°C (986°F). When dissolved in Water and neutralised with Hydrochloric Acid it affords the tests distinctive of Potassium given under that heading. Its aqueous solution possesses a strong alkaline reaction towards red Litmus paper, produces a fine pink tint with Phenolphthalein Solution, and has an alkaline reaction towards Methyl Orange. It is *U.S.P.* required to contain at least 89.7 p.c. of pure Potassium Hydroxide, as determined by titration with Volumetric Sulphuric Acid Solution as indicated below. Few samples approach this figure, although the standard is easy of attainment, the general range found in the author's laboratory is between 78 and 85 p.c. The *U.S.P.* requires that it should contain at least 85 p.c. of pure anhydrous Potassium Hydroxide as determined by titration with Normal Volumetric Sulphuric Acid Solution, using, as shown below, Methyl Orange as an indicator of neutrality. The *P.G.* requires that it shall contain at least 90 p.c. of pure Potassium Hydroxide, as determined by titrating an aliquot portion of a solution containing the *U.S.P.* percent of Potassium Hydroxide with Normal Volumetric Hydrochloric Acid Solution as described below.

The more generally occurring impurities are Arsenic, Copper, Lead, Carbonate, Chloride, Sulphate, and Nitrate. It should not yield any reaction for Arsenic when examined by the modified Gutzeit's test. An aqueous solution slightly acidified with Hydrochloric Acid should

not be darkened in colour nor yield a precipitate with Hydrogen Sulphide Solution. No effervescence should occur on the addition of diluted Sulphuric Acid to a 10 p.c. aqueous solution. Commercial Potash as a rule contains 1 or 2 p.c. of Chloride derived from the Carbonate used in its preparation. The presence of Chloride, Sulphate, and Nitrate may be determined by the Silver Nitrate, Barium Nitrate, and Ferrous Sulphate tests described below. In addition to these impurities it may also contain Aluminium, Calcium, and Silica. When neutralised with Hydrochloric Acid it should yield no white flocculent precipitate on the addition of Ammonia Solution, when the solution is boiled. If the liquid be filtered it should yield no distinct turbidity or a precipitate with Ammonium Oxalate Solution, when dissolved in Water, an excess of Hydrochloric Acid added, and evaporated to dryness, the residue should be completely soluble in Water.

Water or Alcohol (90 p.c.)—An aqueous solution (1–20) should be perfectly clear and colourless, *U.S.P.* 1 gramme dissolved in 2 c.c. of Water, then mixed with 10 c.c. of Alcohol, should leave only a very insignificant residue on standing, *P.G.* 1 gramme dissolved in Water or Alcohol should leave only a trace of sediment, *B.P.*

Sulphuric Acid—10 c.c. of an aqueous solution (1–10) should show no distinct effervescence on the addition of an excess of diluted Sulphuric Acid, *U.S.P.*

Lime Water and Nitric Acid—If a solution of 1 gramme of Potassium Hydroxide in 10 c.c. of Water be boiled with 15 c.c. of Lime Water and filtered, the filtrate with excess of Nitric Acid added to it should not evolve gas bubbles, *P.G.*

Sulphuric Acid and Ferrous Sulphate—If 2 c.c. of a solution (1–20) prepared with diluted Sulphuric Acid be mixed with 2 c.c. of Sulphuric Acid, and 1 c.c. of Ferrous Sulphate T.S. poured over it, no coloured zone should appear, *P.G.*

Barium Nitrate—A solution (1–50) saturated with Nitric Acid should not be immediately affected by T.S. of Barium Nitrate, *P.G.*

Silver Nitrate—A solution (1–50) saturated with Nitric Acid should not become more than opalescent with T.S. of Silver Nitrate, *P.G.*

Time-limit Test—An aqueous solution (1–20) slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *U.S.P.*

Volumetric Determination—16.1 c.c. of Normal Volumetric Sulphuric Acid Solution should be necessary for the neutralisation of a solution of 1 gramme of Caustic Potash in Water or Alcohol (90 p.c.) *B.P.*, 9 c.c. of Normal Volumetric Hydrochloric Acid Solution should be necessary to neutralise 10 c.c. of a solution of 5.6 grammes of Potassium Hydroxide in 100 c.c. of Water, *P.H.*, weigh accurately in a stoppered weighing bottle about 1 gramme of Potassium Hydroxide, dissolve in about 50 c.c. of Water and titrate the solution with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange T.S. as indicator *U.S.P.*

Preparation

LIQUOR POTASSÆ SOLUTION OF POTASH

A clear, colourless or almost colourless, strongly alkaline solution. Liquor Potassæ *B.P.* contains 6.19 p.c. w/v of pure Potassium Hydroxide, corresponding to 6.2 grains in 110 minims or 27 grains in 1 fl. oz. The *U.S.P.* Liquor contains about 5 p.c. w/w of Potassium

Hydroxide, corresponding to 5.23 p c w/v, about $5\frac{1}{4}$ grains per 110 minims or 23 grains per fl oz. The *P G* Liquor contains 15 p c w/w of pure Potassium Hydroxide, equivalent to 5.23 p c w/v, 17.1 grains per 110 minims or 75 grains per fl oz.

It should be preserved in well-stoppered glass bottles of a dark amber shade, and the stoppers may be smeared with mineral Oil to prevent fixation.

Medicinal Properties.—Caustic. When diluted it is antacid and antispasmodic. Occasionally employed as an antacid in dyspepsia, accompanied by acidity and gastralgia. It is apt to irritate the stomach, and so, to obtain all the best internal effects of Potash, the Bicarbonate and Citrate are much to be preferred. Externally as an escharotic against the bite of rabid or venomous animals, diluted, it relieves itching.

It acts powerfully on all organic matter, converting flannel into a kind of soft jelly after immersion for 5 or 6 hours.

Dose—10 to 30 minims = 0.6 to 1.8 c c, freely diluted.

Incompatibles—Acids, acid salts, metallic and alkaloidal salts, the preparations of Ammonium, Belladonna, Henbane, and Stramonium.

Antidotes—Diluted Acetic Acid, Citric Acid, Lemon Juice, or any vegetable acids, fixed oils and demulcents, stimulants, Morphine for pain, neither stomach-tube nor emetics are to be used.

Foreign Pharmacopœias—Official in U.S., sp gr about 1.046 at 25° C (77° F) (5 p c), Austr (Kalium Hydroxydatum Solutum), sp gr 1.325 to 1.332, Dutch (Solutio Hydratis Kalici), sp gr 1.180, Fr (Hydroxyde de Potassium dissous), sp gr 1.080, Ger (Liquor Kali Caustici), sp gr 1.138 to 1.140 (15 p c), Russ (Kali Causticum Solutum), sp gr 1.126 to 1.130 (15 p c), Swed (Solutio Hydratis Kalici), sp gr 1.225 to 1.235 (25 p c), Swiss (Kalium Hydricum Solutum), sp gr 1.33. Not in the others.

Tests—Liquor Potassæ is officially required to have a sp gr of 1.058. The *USP* gravity is 1.046 at 25° C (77° F). The *P G* 1.138 to 1.140. It possesses a strong alkaline reaction towards blue Litmus paper and towards Phenolphthalein and Methyl Orange Solutions. It answers the tests distinctive of Potassium given under that heading. It is officially required to contain 6.19 p c w/v, equivalent to 5.85 p c w/w of pure Potassium Hydroxide as volumetrically determined by titration with Volumetric Sulphuric Acid Solution as described below. The *USP* requires that it shall contain about 5 p c w/w, equivalent to 5.23 p c w/v of Potassium Hydroxide as volumetrically determined by the process given below, using Methyl Orange Solution as an indicator of neutrality. The *P G* requires that it shall contain about 15 p c w/w, equivalent to 17.1 p c w/v of Potassium Hydroxide, but does not indicate a method by which this requisite percentage can be assured. The Liquor should be free from the more generally mentioned impurities mentioned under Potassa Caustica. It is also required to be free from Ammonia, Magnesium, Iron or Sodium. Why Potassium Hydroxide should be required to yield no characteristic reaction with the tests for Arsenic, Copper and Lead only, whilst the solution made therefrom should be required not to yield any characteristic reaction with

the tests for Aluminium, Ammonium, Arsenic, Calcium, Copper, Iron, Lead, Magnesium and Sodium, as well as to be free from more than traces of Carbonates, Chlorides or Sulphates, is not quite apparent. The *USP* more consistently remarks that it should conform to the reactions and tests for an aqueous Potassium Hydroxide Solution given under Potassu Hydroxidum. The aqueous solution should not evolve ammoniacal odours when boiled, nor should the issuing vapour have an alkaline reaction towards red Litmus paper. The aqueous solution acidified with Hydrochloric Acid treated with Ammonia in slight excess, and boiled and filtered, treated with Ammonium Oxalate Solution and again filtered, should yield no distinct turbidity nor a precipitate on the addition of Sodium Phosphate Solution. When freshly made, Potassium Hydroxide Solution usually contains a little Lime, but as it absorbs Carbon Dioxide the Lime is thrown out.

Lime Water and Nitric Acid—The solution, when boiled with 4 parts of Lime Water and filtered, gives a filtrate which does not evolve gas bubbles with excess of Nitric Acid, *P G*.

Barium Nitrate—Diluted with 5 parts of Water and supersaturated with Nitric Acid, it should not become more than opalescent with T S of Barium Nitrate, *P G*.

Silver Nitrate—A similar solution should not become more than opalescent with T S of Silver Nitrate, *P G*.

Sulphuric Acid and Ferrous Sulphate—If 2 c c of Potassium Hydroxide be neutralised with diluted Sulphuric Acid, then mixed with 2 c c of Sulphuric Acid and 1 c c of Ferrous Sulphate T S poured on as a layer, no coloured zone should be produced, *P G*.

Ammonia—After supersaturating solution of Potassium Hydroxide with Hydrochloric Acid, it should not become more than opalescent with Ammonia T S even after standing, *P G*.

Volumetric Determination—10 c c of Normal Volumetric Sulphuric Acid Solution neutralises 9 c c of Solution of Potash, *BP*, 25 c c of Normal Volumetric Sulphuric Acid Solution should be necessary to neutralise 28 (27.87) grammes of solution of Potassium Hydroxide, using Methyl Orange T S as indicator, *USP*.

Not Official

BRANDISH'S ALKALINE SOLUTION—American Pearl ash, 6 lbs, freshly prepared Quicksilver, 2 lbs, Wood ashes, 2 lbs, boiling Water, 6 gallons or 6, 2, 2, and 60 parts, add first the Lime, then the Pearl ash, and lastly the Wood ashes to the boiling Water, stir well together, let it stand 24 hours and decant the clear liquor.

Dose— $\frac{1}{2}$ to 2 fl drms = 1.8 to 7.1 c c in Milk. Given for scrofulous conditions.

POTASSA CUM CALCE (Vienna Paste)—Potassium Hydroxide and Calcium Oxide, equal weights, powder and mix, it is made into a paste with Alcohol (90 p c).

This has been incorporated in the *BP C*.

The paste is spread on the part to be cauterised, and is allowed to remain for 10 to 15 minutes, while the surrounding skin is protected by adhesive plaster. It is also used in the treatment of lupus. It is not so likely to diffuse as Caustic Potash alone.

Foreign Pharmacopœias—Official in Fr and Ital, Potassium Hydroxide 5, Lime 6, Mex (Pasta de Viena), Potassium Hydroxide 1, Lime 1, Span. (Caustico de Viena), Potassium Hydroxide 50, Lime 60.

Potassa cum Calce in cylinders, consisting of 2 parts of Potassa and 1 of Lime for the use of gynecologists.

POTASSA SULPHURATA.

SULPHURATED POTASH

B P Syn—LIVER OF SULPHUR

N O Syn—HEPAR SULPHURIS

FR, SULFURE DE POTASSE, GER, SCHWEFFELKREDER, ITAL, SOLFURO DI POTASSIO, SPAN, SULFURO (TRI) POTASICO

Liver-brown, deliquescent, irregular pieces, which gradually absorb moisture and Carbon Dioxide, the colour changing to greenish-yellow. It has a " " odour of Hydrogen Sulphide when slightly moist, and an alkaline reaction. It is a mixture of various Potassium salts, chiefly Sulphides.

As it is deliquescent and liable to oxidation on exposure to the air it should be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from contact with the air, it should also be kept in a cool atmosphere.

Solubility—1 in 2 of Water

Medicinal Properties.—Similar to those of Sulphur, but more energetic. Externally, as a bath or in ointment form, it is a good remedy for scabies and other parasitic cutaneous diseases, used also for chronic eruptions, " " psoriasis and acne. Internally it is occasionally used for chronic rheumatism and chronic skin diseases.

A hot bath of Sulphurated Potash relieves the itching of jaundice—*L* '85, ii. 1220

Dose—1 to 5 grains = 0.06 to 0.32 gramme

Not Official—Unguentum Potassæ Sulphuratæ, Pommade Sulfureuse, Balneum Sulphuretum, Bain Sulfuré, Bain Sulfuré Liquide

Foreign Pharmacopœias—Official in Austr, Ger, Jap, Russ and Swiss (Kalium Sulfuratum), Austr and Hung have (Kalium Sulfuratum pro Balneo), Belg (Sulphuretum Potassii Officinale), Dan, Norw, and Swed (Hepar Sulphuris), Dutch (Trisulphuretum Kalicum), Fr (Sulfure de Potasse), Ital (Solfuro di Potassio), Mex (Sulfuro de Potasio), Port (Potassa Sulfurada), Span (Sulfuro (tri) Potasico) Not in U.S.

Tests—Potassium Sulphide dissolves readily in Water, forming a yellow solution possessing an odour of Hydrogen Sulphide and an alkaline reaction towards red Litmus paper. On the addition of an excess of Hydrochloric Acid it evolves the distinctive odour of Hydrogen Sulphide, and the issuing gas produces a black stain on paper moistened with Lead Acetate Solution, a deposit of Sulphur simultaneously appears in the liquid. When freed from Hydrogen Sulphide by boiling until the vapours no longer cause a discoloration on Lead Acetate paper, and filtering, the filtrate yields the tests distinctive of Potassium given under that heading. A portion of the filtrate also yields with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. The *B P* requires that about 50 p.c. of the substance should be soluble in Alcohol (90 p.c.) When well made it contains from 50 to 60 p.c. of Potassium Sulphide.

Not Official

UNGUENTUM POTASSÆ SULPHURATÆ—Sulphurated Potash, 30 grains, Hard Paraffin, $\frac{1}{4}$ oz, Soft Paraffin, $\frac{3}{4}$ oz This ointment should be freshly prepared —*B P* 1885

This has been incorporated in the *B P C*

POMMADE SULFUREUSE (*Vet*) —Sulphurated Potash (powdered very finely), 1, Vaseline, 30 Mix to form a homogeneous pomade —*M*

BALNEUM SULPHURETUM—Sulphurated Potash, 4 oz, Water, 30 gallons, dissolve

Used in psoriasis, etc

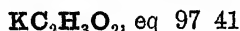
This is not quite so agreeable as the Bâleges waters, which may be made artificially as follows —Sodium Sulphide, Sodium Carbonate, and Sodium Chloride, of each 20 grains to 1 gallon But a much stronger solution is often used

BAIN SULFURÉ—100 grammes of Sulphurated Potash (broken up) are placed in well closed bottle When required for use, it is dissolved in a litre of warm water and the solution is poured into the water bath The bath should be non metallic or made of zinc —*F*

BAIN SULFURÉ LIQUIDE—Sulphurated Potash, 1, Water, 2 Dissolve, filter and preserve in a well-closed bottle to be added to a bath when required —*F*

POTASSII ACETAS.

POTASSIUM ACETATE



FR, ACETATE DE POTASSIUM, GER, KALIUMACETAT, ITAL, ACETATO DI POTASSIO, SPAN, ACETATO POTÁSICO

White, almost odourless, deliquescent crystals, or crystalline satin-like masses, or as a white, deliquescent, coarse granular powder

It should be kept in well closed vessels and protected as far as possible from contact with the air

It may be prepared by neutralising Potassium Carbonate with Acetic Acid, evaporating and fusing the product It has been stated that the formula given in the *B P* does not represent the substance actually in use, which cannot be rendered anhydrous on a technical scale without decomposition An allowance of 10 p c of Water on drying at 110° C (230° F) would be reasonable

Solubility—2 in 1 of Water, 1 in 2 of Alcohol (90 p c)

Medicinal Properties—Used as a diuretic in dropsy, chiefly renal, and in febrile diseases, as an antilithic in gout and the uric acid diathesis, valuable in sub-acute rheumatism

Dose.—10 to 60 grains = 0.65 to 4 grammes

Prescribing Notes—*Best administered in simple solution, with a little Syrup if necessary*

Foreign Pharmacopœias—Official in all except Austr, Ger, Jap and Swiss, Austr, contains a *solution*, sp gr 1.200, Ger, Hung, Ital, Jap and Russ have also a solution, sp gr 1.176 to 1.180 (about 83 p c), Swiss has Liquor, sp gr 1.17 to 1.18

Tests.—Potassium Acetate fuses when strongly heated, the *U S P* states at a temperature of 292°C (557.6°F). At a still higher temperature it chars, and when ignited it should leave a white residue completely soluble in Water. The salt dissolves readily in Water, forming a clear solution which is alkaline in reaction towards red Litmus paper, but which does not produce a pink coloration with Phenolphthalein Solution. It yields the tests distinctive of Potassium given under that heading. The aqueous solution when mixed with Sulphuric Acid and boiled evolves a distinctive acetous odour. When warmed with Sulphuric Acid and a few drops of Alcohol (90 p c) a distinctive odour of Ethyl Acetate is evolved. An aqueous solution yields with Ferric Chloride T S a deep red coloration, and on boiling a brown flocculent precipitate is thrown out. The *BP* does not require it to contain any definite of pure Potassium Acetate, nor is a method given by an be assayed. The *U S P* requires that it should contain when thoroughly dry not less than 98 p c of pure Potassium Acetate as volumetrically determined by titration of the solution of the residue left on ignition with Semi-normal Volumetric Sulphuric Acid Solution as indicated below, using Methyl Orange Solution as an indicator of neutrality.

The more generally occurring impurities are Aluminium, Arsenic, Calcium, Copper, Iron, Lead and Magnesium, the *BP* requiring that the usual 'no characteristic reaction' should be yielded with the tests for these impurities, and also for Carbonates or Sulphides. Chlorides and Sulphates may also be present, the official directions are that only the slightest reactions with the tests for these substances shall be yielded. Arsenic, if present, may be detected by the alliaceous odour evolved during the cautious ignition of the sample. It may also be determined by the Hydrogen Sulphide test given below, and by the Gutzzeit's test also described below. Copper, Iron and Lead, if present, may be recognised by the Hydrogen Sulphide test either in acid or alkaline solution. No flocculent precipitate nor a turbidity should be produced by the addition of Ammonia Solution on boiling. The filtrate should not yield a turbidity nor a precipitate with Ammonium Oxalate Solution. If the liquid be again filtered, the filtrate should neither yield a turbidity nor a precipitate with Sodium Phosphate Solution. Chlorides and Sulphates, if present, are indicated by Silver Nitrate and Barium Chloride Solutions.

Time-limit Test.—An aqueous solution (1-20) slightly acidulated with Acetic Acid should not respond to the time-limit test for heavy metals, *U S P*.

Gutzzeit's Test.—5 c c of an aqueous solution of the salt (1-10) should not respond to the modified Gutzzeit's test for Arsenic, *U S P*.

Volumetric Determination.—Thoroughly carbonise 1 gramme of dry Potassium Acetate at a temperature not exceeding red heat, extract the residue with boiling Distilled Water until the washings cease to react with Methyl Orange T S. The mixed filtrate and washings should require for complete neutralisation not less than 20.1 c c of Semi-normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as indicator, *U S P*.

Not Official

POTASSII BENZOAS

 $\text{KC}_7\text{H}_5\text{O} \cdot 3\text{H}_2\text{O}$, eq 212.60

A white crystalline powder, soluble 1 in $1\frac{1}{2}$ of Water, 1 in 18 of Alcohol (90 p c). Useful in cystitis of gouty or rheumatic origin.

It should be kept in well stoppered glass bottles and in a cool atmosphere.

Dose—15 to 20 grains = 1 to 1.3 grammes

Tests—Potassium Benzoate dissolves readily in Water, yielding a solution which is slightly alkaline in reaction towards red Litmus paper, and which affords, if sufficiently concentrated, on the addition of diluted Sulphuric Acid Solution, a copious white crystalline precipitate. If this precipitate be separated, washed and carefully dried, it should possess the m p and answer the tests distinctive of Benzoic Acid given under Acidum Benzoicum. The filtrate from the precipitate yields the tests distinctive of Potassium given under that heading. An aqueous solution of the salt affords, with Ferric Chloride T.S., a buff coloured precipitate. The percentage of pure Potassium Benzoate may be determined by dissolving 1 gramme in Water, adding sufficient Ether to dissolve the Benzoic Acid and titrating with Tenth normal Volumetric Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutrality. 1 c.c. of Tenth-normal Volumetric Hydrochloric Acid Solution corresponds to 0.02126 gramme of pure Potassium Benzoate.

It may also be assayed by igniting the Benzoate and titrating the filtered solution of the residual Potassium Carbonate with Tenth normal Volumetric Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutrality.

POTASSII BICARBONAS.

POTASSIUM BICARBONATE

B. P. Syn—POTASSIUM HYDROGEN CARBONATE KHCO_3 , eq 99.38

FR., CARBONATE ACIDE DE POTASSIUM, GER., KALIUMBICARBONAT,
ITAL., BICARBONATO DI POTASSIO, SPAN., BICARBONATO POTASICO

Colourless, transparent, monoclinic prisms, or as a white crystalline powder, odourless and having a saline and slightly alkaline taste. It is permanent in the air.

It may be obtained by saturating a strong aqueous Potassium Carbonate solution with Carbonic Anhydride.

It should be kept in well-closed vessels and in a cool atmosphere.

Solubility—1 in 3.2 of Water. Insoluble in Alcohol (90 p c).

Medicinal Properties—Antacid, antilithic, and diuretic. Used in dyspepsia as an antacid, and in gout to increase the alkalinity of the blood and excretion of urates, in the acute or inflammatory stage of gonorrhoea there is no better remedy, as it renders the urine alkaline and unirritating. In bronchitis and pneumonia it renders the secretion less tenacious, in influenza it has been given with success. See also Sodium Bicarbonate, which is generally preferred in dyspepsia.

20 grains are prescribed in effervescence with 15 grains of Citric Acid.

Closely resembles the Carbonate, but without its irritant qualities.

Potassium salts delay the conversion of gelatinous Sodium Biurate into the

crystalline variety, and when the conversion is once started it is slowed by the presence by these salts. Potassium salts exercise most influence.—*L* '00, 1 931

Given in the treatment of gleet in order to artificially produce phosphaturia.—*L* '08, 1 424

Dose.—5 to 30 grains = 0.32 to 2 grammes

Foreign Pharmacopœias.—Official in U.S., Belg. (Bicarbonas Potassæ), Fr. (Carbonate Acide de Potassium), Norw. and Swed. (Bicarbonas Kalicus), Ger., Jap., Russ. and Swiss (Kalium Bicarbonicum), Mex. (Carbonato de Potasio ácido), Port. (Bicarbonato de Potassa), Span. (Bicarbonato Potásico). Not in Austri., Dan., Dutch, Hung., or Ital.

20 parts by weight of Potassium Bicarbonate are neutralised by 14 parts of Citric Acid, and by 15 parts of Tartaric Acid.

Tests.—Potassium Bicarbonate loses Carbonic Anhydride when exposed to a temperature of about 100° C (212° F) and at a dull red heat is completely converted into Potassium Carbonate. It dissolves in Water, forming a clear solution which is alkaline in reaction towards red Litmus paper, but neutral in reaction towards Phenolphthalein Solution. It answers the tests distinctive of Potassium given under that heading, it dissolves with effervescence in diluted Sulphuric Acid, the evolved gas yielding, when passed into Lime Water, a white precipitate soluble in a sufficient excess of gas. It is officially required to contain 99.4 p.c. of pure Potassium Bicarbonate as determined, first, gravimetrically by the weight of residue left on ignition, and, secondly, volumetrically by the titration of that residue with Volumetric Sulphuric Acid Solution as described below. The *U.S.P.* requires that it shall contain not less than 99 p.c. of pure Potassium Bicarbonate as volumetrically determined by direct titration with Semi-normal Volumetric Sulphuric Acid Solution, Methyl Orange T.S. being employed as an indicator of neutrality. The *P.G.* requires that it shall contain 100 p.c. of pure Potassium Bicarbonate, first, as volumetrically determined by direct titration with Normal Volumetric Hydrochloric Acid Solution, secondly, as gravimetrically determined by the weight of residue left on ignition. Both the *U.S.P.* volumetric determination method and the volumetric and gravimetric methods of the *P.G.* are described in the small type below.

The following are the impurities occurring: Arsenic, Calcium, Copper, Iron and Lead, Carbonates, Chlorides, Nitrates and Sulphates. The *B.P.* requires that in addition to these it shall yield no characteristic reaction with the tests for Aluminium, Magnesium and Sodium and Sulphides. Arsenic, if present, may be detected by the modified Gutzeit's test, using the Hydrochloric Acid solution of the salt, and also by the Hydrogen Sulphide test described below, which also detects, if present, Copper and Lead. A standard has been suggested (*C.D.* '08, 1 796) of 5 parts per million for Lead and 2 parts per million for Arsenic, and for Chlorides a standard of 0.1 p.c. calculated as KCl. The *P.G.* includes a separate test for Iron with Potassium Ferrocyanide, see below. Calcium, if present, may be detected by a turbidity or precipitate produced on adding Ammonium Oxalate Solution to an aqueous solution of the salt rendered slightly acid with Acetic Acid. Magnesium, if present, may be detected in the

filtrate from the Ammonium Oxalate Solution Aluminium and Sodium are unlikely impurities Chlorides and Sulphates, if present, may be detected by the Barium Nitrate and Silver Nitrate test described below Nitrates may be detected by the Ferrous Sulphate and Sulphuric Acid test The *USP* uses the Phenolphthalein test described below to detect the presence of Carbonate

Phenolphthalein.—A concentrated aqueous solution is neutral to T S of Phenolphthalein, *USP* Dissolve 1 gramme of the salt without agitation in 20 c c of Water at a temperature not above 15° C (59° F), add 0.2 c c of Normal Volumetric Hydrochloric Acid Solution and 2 drops of Phenolphthalein T S A red tint should not appear immediately, *USP*

Barium Nitrate—An aqueous solution (1-20) after saturation with Acetic Acid should not be affected by T S of Barium Nitrate, *PG*

Silver Nitrate—A solution as above, after the addition of Nitric Acid, should not become more than opalescent with T S of Silver Nitrate, *PG*

Hydrogen Sulphide—An aqueous solution (1-20) after saturation with Acetic Acid should not be affected by T S of Hydrogen Sulphide, *PG*, a (1-20) aqueous solution slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*

Potassium Ferrocyanide—20 c c of an aqueous solution (1-20) saturated with Hydrochloric Acid should not become blue with 0.5 c c of T S of Potassium Ferrocyanide, *PG*

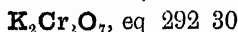
Gravimetric Determination—1 gramme when heated at a low red heat leaves 0.69 gramme of a white residue, *BP*, 100 parts by weight of the salt when ignited to a dull red heat shall leave 69 parts by weight of residue, *PG*

Volumetric Determination—The 0.69 gramme of white residue obtained by igniting 1 gramme of the salt at a dull red heat shall require 10 c c of Volumetric Sulphuric Acid Solution for neutralisation, *BP* 1 gramme should require for neutralisation 10 c c of Normal Volumetric Solution of Hydrochloric Acid, *PG* 1 gramme should require for neutralisation not less than 19.9 c c of Semi normal Volumetric Solution of Sulphuric Acid, using Methyl Orange as indicator, *USP*

POTASSII BICHROMAS.

POTASSIUM BICHROMATE

BP Syn—POTASSIUM DICHROMATE, RED CHROMATE OF POTASSIUM



FR, CHROMATE ACIDE DE POTASSIUM, GER, KALIUMDICHROMAT, ITAL, BICROMATO DI POTASSIO, SPAN, BICROMATO POTASICO

Large, orange-red, odourless, translucent, prismatic crystals, having a bitter taste Permanent in the air

It should be kept in well stoppered glass bottles of a dark amber tint in a cool place

Solubility—1 in 10 of Water, 5 in 6 of boiling Water

Medicinal Properties—A powerful irritant poison in overdoses, rarely used in medicine, but extensively in the arts

Highly recommended by Fraser in dyspepsia and gastric ulcer (*L* '94, 1 923), and by Bradbury—*L* '95, 11 671

Dose— $\frac{1}{10}$ to $\frac{1}{5}$ of a grain = 0.006 to 0.013 gramme, in pills with 'Massa Kaolin'

Official Preparation—Used in the preparation of Acidum Chromicum

Antidotes.—Stomach-tube or emetics, Magnesium Carbonate or Chalk, albuminous and demulcent drinks

Foreign Pharmacopœias—Official in Fr, Ger, Port, Russ, Span, Swed, Swiss and U S Not in the others

Tests—Potassium Bichromate fuses below a red heat, and at a considerably higher temperature is decomposed, evolving Oxygen and leaving a residue of neutral yellow Chromate and green Chromium Oxide. The salt dissolves in Water with the formation of a reddish-yellow solution, which has a slightly acid reaction towards blue Litmus paper. When warmed with Sulphuric Acid and Ethylic Alcohol the yellow aqueous solution assumes a green colour, at the same time evolving a distinctive odour of Acetaldehyde. The experiment requires to be carefully conducted, as the reaction is very energetic. An aqueous solution yields, with Barium Chloride, a reddish precipitate soluble in Hydrochloric Acid. With Silver Nitrate Solution it yields a dark purple-red precipitate soluble in Nitric Acid and in Ammonia Solution. If sufficient Barium Chloride Solution or Silver Nitrate Solution be added to an aqueous solution of the salt to precipitate the whole of the Chromium as Chromate, the filtrate from these precipitates answers the test distinctive of Potassium given under that heading. It is officially required to contain 99.8 p.c. of pure Potassium Bichromate as volumetrically determined by the process described in the small type below. The *USP* requires that it should contain not less than 99 p.c. of pure Potassium Bichromate, but does not give a method of determination by which this requisite percentage may be assured. The *PG* gives neither a percentage nor a method of determination.

The more generally occurring impurities are Calcium, Chlorides and Sulphates. Chlorides and Sulphates, if present, may be detected by the Barium Chloride and Silver Nitrate tests in acid solution, Calcium, if present, by the Ammonium Oxalate test described below.

Barium Chloride or Nitrate—The yellowish-white precipitate obtained on adding Barium Chloride T.S. to an aqueous solution of the salt should be entirely soluble in diluted Nitric Acid, *BP*, the *PG* states that an aqueous solution (1-100) strongly acidulated with Nitric Acid and warmed should not be affected by Barium Nitrate T.S.

Silver Nitrate—The dark purple-red precipitate obtained on adding Silver Nitrate to an aqueous solution of the salt should be entirely soluble in Nitric Acid, *BP*, the *PG* states that an aqueous solution as in the previous test should not be affected by T.S. of Silver Nitrate.

Ammonium Oxalate—An aqueous solution (1-100) of the salt after the addition of Ammonia T.S. should not become turbid with T.S. of Ammonium Oxalate, *PG*.

Volumetric Determination.—To effect the oxidation of the Iron from the Ferrous to the Ferric condition a solution of 5.66 grammes of Ferrous Sulphate in Water acidulated with Sulphuric Acid, *BP*, to which the addition of a solution containing 1 gramme of Potassium Bichromate, *BP*.

POTASSII BROMIDUM.

POTASSIUM BROMIDE

NO Syn—BROMURETUM KALICI**KBr**, eq 118 18

Fr, BROMURE DE POTASSIUM, Ger, KALIUMBROMID, Ital, BROMURO DI POTASSIO, Span, BROMURO POTASICO

Colourless or white, odourless, cubical crystals, possessing a strong, characteristic, saline taste They are permanent in the air

It should be kept in well-closed bottles and protected as far as possible from the light

Solubility—10 in 17 of Water, and measures 20, 1 in 1 of boiling Water, 1 in 95 of Alcohol (90 p c), 1 in 17 of boiling Alcohol (90 p c)

Medicinal Properties—Sedative, hypnotic, anaphrodisiac Very successful in epilepsy, in hysteria, and in convulsions generally Used in insomnia, due not to pain but to worry or overwork or the climacteric, sea-sickness and the sickness of pregnancy, also in nervous headache, nightmare and the night-screaming of children, in migraine and in neuralgia Useful in spermatorrhoea and nymphomania, and with chloral in delirium tremens Relieves in some cases of whooping-cough and spasmodic asthma, both in children and adults In enormous doses sometimes successful in tetanus

Bromides still the most potent agents for the treatment of idiopathic epilepsy No special advantage noticed in giving the mixed Potassium, Sodium and Ammonium Bromides Potassium salt usually given, but when it appears to cause undue depression, the Ammonium or Sodium salt is substituted In nocturnal epilepsy, a single dose should be given an hour before bed time, and two hours before an attack is due when the fits recur about the same time in the day As a rule, better to increase the single dose than to give the same amount in divided doses several times a day—*L* '08, 1 440, *B M J* '08, 1 871

Bromide rash treated by use of arsenical waters, milk diet and a poultice containing Boric Acid applied locally—*T G* '99, 598

On its use combined with Sodium Salicylate in headache—(Brunton) *Pr* 101

By combining it with Arsenic in small doses, the unpleasant effects known as 'Bromism' may be prevented or reduced

Butter milk as a local application to the acne like eruption produced by administration of Bromides—*L* '02, 11 1724

Dose—5 to 30 grains = 0.32 to 2 grammes

Incompatibles—Any oxidising agents are liable to set free the Bromine, Spiritus Ætheris Nitrosi

Official Preparation—Used in the preparation of Acidum Hydrobromicum Dilutum

Not Official—Sal Bromatum Efflorescens

Foreign Pharmacopœias—Official in Austr, Ger, Hung, Jap, Russ and Swiss (Kalium Bromatum), Belg (Bromuretum Potassii), Dan, Dutch, Norw and Swed (Brometum Kalicum), Fr (Bromure de Potassium), Ital (Bromuro di Potassio), Mex (Bromuro de Potasio), Port (Brometo de Potassio), Span (Bromuro Potasico), US (Potassii Bromidum)

Tests—Potassium Bromide when heated decrepitates and when strongly heated fuses. It dissolves readily in Water, forming a clear solution which should be neutral in reaction towards Litmus paper. It answers the tests distinctive of Potassium given under that heading. The aqueous solution, when treated with a little Chlorine Water, assumes a brown or reddish-brown coloration, and when shaken with Chloroform the colour passes into the chloroformic solution. On the addition of Silver Nitrate Solution the aqueous solution affords a yellowish, curdy precipitate, practically insoluble in Ammonia Solution, insoluble in Nitric Acid, readily soluble in Potassium Cyanide Solution. A small quantity of the salt heated with Manganese Dioxide and Sulphuric Acid evolves the characteristic irritating vapours of Bromine, which communicates an orange-yellow colour to filter paper soaked in Starch Mucilage. It is officially required to yield not less than 98.9 p.c. nor more than 100.9 p.c. of pure Potassium Bromide as volumetrically determined by direct titration of the salt with Volumetric Silver Nitrate Solution as indicated below. The *U.S.P.* requires that it should contain not less than 97 p.c. of pure Potassium Bromide as volumetrically determined by direct titration of the well-dried salt with Tenth-normal Volumetric Silver Nitrate Solution as described below, using Potassium Chromate Solution as an indicator. The *P.G.* requires that it shall contain not more than 100.8 p.c. of pure Potassium Bromide as determined by titrating an aliquot portion of a solution of 3 grammes of the salt dried at 100° C (212° F) dissolved in 100 c.c. of Water, as described in small type below, using Potassium Chromate Solution as an indicator. If the figures required by the official Silver Nitrate titration be calculated into *KBr* they would indicate a percentage of 98.91 to 100.92. As 100 p.c. *KBr* requires 84.62 c.c., the excess over the theoretical figure would be due to *KCl*, which may be present from 0.1 p.c. to 6 p.c. This cannot give a definite Chloride figure unless all impurities unaffected by Silver Nitrate are known to be absent, the only interfering impurity, however, which may be expected to be present is Water, so that if *B.P.* had directed the dried salt to be used for titration, the percentage of Chloride might be arrived at by $\frac{84.62}{0.5}$ from the number of c.c. used and dividing the result by 0.5. Some English samples of the salt contain less than $\frac{1}{4}$ p.c. of Chloride, but some American samples contain nearly 6 p.c.

The more generally occurring impurities are Arsenic, Copper, Iron, Lead and Zinc, Barium and Calcium, Bromates, Iodides and Iodates, Chlorides and Sulphates. In addition to the usual official statement that it should yield 'no characteristic reaction' for the tests for Aluminium, or only the slightest reaction with the tests for these substances, the *B.P.* also includes similar requirements with regard to Ammonium, Magnesium, Sodium, and gives a specific and definite test for absence of Thiocyanates. Arsenic, if present, may be detected by the Gutzzeit's test. It may also, together with Copper, Iron, Lead and Zinc, if present, be detected by the Hydrogen Sulphide test described below, Arsenic, Copper and Lead

in slightly acid solution, Iron and Zinc in alkaline solution Barium, if present, may be detected by the Potassium Sulphate test described below Calcium, if present, by the addition of Ammonium Oxalate to the aqueous solution slightly acidified with Acetic Acid Bromates, if present, may be detected by the Sulphuric Acid test given below The Barium Nitrate test serves to detect Sulphates if present The Chlorine Water and Chloroform test described below serves to detect the presence of Iodides Any excessive proportion of Chloride may be detected by the increase in the titration figure as indicated above The *PG* includes a separate test for Iron with Potassium Ferrocyanide Solution The *USP* includes a test for limit of alkali, which is given under the Phenolphthalein test in small type below

The *BP* requires that the cold aqueous solution should not assume a red coloration on the addition of Ferric Chloride TS It has been suggested (*Pharm. J.* '01, 1 460) that this Thiocyanate test requires modification, the colour produced by Ferric Chloride in an aqueous solution of Potassium Bromide largely depending on the quantity of Potassium Bromide present, the colour being masked by an excess of Potassium Bromide The modification proposed is that 2 drops of Ferric Chloride TS should give a yellow, and not a red or reddish-brown, coloration when added to a solution of 0.1 gramme of Potassium Bromide dissolved in 10 c.c. of Water The test is stated to indicate the absence of more than 0.1 per cent of Ammonium Thiocyanate

Litmus—An aqueous solution (1-20) is neutral or has only a scarcely perceptible reaction on Litmus, *USP* Powdered Potassium Bromide should not immediately colour moistened red Litmus paper violet blue, *PG*

Phenolphthalein—A solution of 1 gramme of the salt in 10 c.c. of Water with 0.1 c.c. Tenth normal Volumetric Sulphuric Acid Solution added should not give any coloration on the subsequent addition of a drop of TS of Phenolphthalein, *USP*

Diluted Sulphuric Acid—Crushed Potassium Bromide spread out on white porcelain should not immediately turn yellow on the addition of diluted Sulphuric Acid, *PG* Such a mixture, when shaken with 1 c.c. of Chloroform, should not impart to the latter a yellowish brown colour, *USP*

Hydrogen Sulphide—An aqueous solution (1-20) of the salt should not be affected by TS of Hydrogen Sulphide, *PG* A solution of similar strength acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*

Barium Nitrate—An aqueous solution of the salt (1-20) should not be affected by TS of Barium Nitrate, *PG*

Potassium Sulphate or Diluted Sulphuric Acid—An aqueous solution of the salt (1-20) should not be affected by diluted Sulphuric Acid, *PG*, 10 c.c. of an aqueous solution (1-20) acidulated with Hydrochloric Acid should not be rendered turbid by the addition of 1 c.c. of TS of Potassium Sulphate, *USP*

Potassium Ferrocyanide—20 c.c. of an aqueous solution (1-20) should not be rendered blue by 0.5 c.c. of TS of Potassium Ferrocyanide, *PG*

Diluted Chlorine Water and Chloroform—Add 1 c.c. of Chloroform to 10 c.c. of an aqueous solution of the salt (1-20) and then introduce cautiously drop by drop with constant agitation a mixture of equal volumes of Chlorine Water and Water The liberated Bromine will dissolve in the Chloroform, imparting to it a yellow to orange colour free from any violet tint, *USP*

Volumetric Determination—A solution of 1 gramme of the salt in Water should require for complete precipitation not less than 88.7 nor more than 85.4 c.c. of Volumetric Solution of Silver Nitrate, *BP*, 10 c.c. of an (3 grammes in 100 c.c.) of Potassium Bromide which has been (212° F) should after the addition of a few drops of Potassium Chromate T.S. require not more than 25.4 c.c. of Tenth-normal Volumetric Solution of Silver Nitrate to produce a permanent reddening, *PG*, 0.3 gramme of the well dried salt dissolved in about 50 c.c. of Water and 2 or 3 drops of Potassium Chromate T.S. added should require not less than 24.6 c.c. nor more than 25.85 c.c. of Tenth-normal Silver Nitrate Volumetric Solution to produce a permanent red colour, *USP*.

Not Official

SAL. BROMATUM EFFERVESCENS—An effervescent preparation containing about 8 of Potassium Bromide, 8 of Sodium Bromide and 4 of Ammonium Bromide in 60.

CALCIUM BROMIDE—A white, very deliquescent salt, readily soluble in Water and Alcohol (90 p.c.), which has been introduced as a substitute for Potassium Bromide, but which is not much prescribed.

Dose—15 to 30 grains = 1 to 2 grammes

Official in US

POTASSII CARBONAS.

POTASSIUM CARBONATE

BP Syn—SALT OF TARTAR

FR, CARBONATE NEUTRE DE POTASSIUM, GER, KALIUMCARBONAT, ITAL, CARBONATO DI POTASSIO, SPAN, CARBONATO POTASICO

A white, deliquescent, crystalline, or granular powder, K_2CO_3 , eq 137.21, officially stated to be associated with either 1 or 2 molecules of Water. It is present in the ashes of plants.

Should be preserved in well-closed bottles.

The *BP* Carbonate is associated with either 1 or 2 molecules of Water of crystallisation. The *USP* Carbonate is required to contain when thoroughly dry not less than 98 p.c. of pure Potassium Carbonate. The *PG* requires it to contain at least 95 p.c. of pure Potassium Carbonate.

It has been stated that the article as met with in commerce is not a definite hydrate, but with 1 or 2 molecules of Water, but a mixture of combined Water. Six samples obtained from leading manufacturers examined in the author's laboratory lost from 16.4 to 19.8 p.c. after exposure to a red heat, the loss averaged 17.4 p.c. The percentage of Chloride present varied between 0.052 and 0.08 p.c., with an average of 0.123 p.c.

Solubility—4 in 3 of Water, and measures 4½. Insoluble in Absolute Alcohol.

Medicinal Properties.—Similar to those of the Bicarbonate, but rarely used internally on account of its irritant properties. Externally it is used as a lotion in eczema and urticaria.

Dose—5 to 20 grains = 0.32 to 1.3 gramme

Official Preparations—Contained in Decoctum Aloes Compositum, Liquor Arsenicalis, Mistura Ferri Composita, Unguentum Potassii Iodici, in the preparation of Iodoform, Liquor Bismuthi et Ammonii Citratis, Potassa Caustica, Potassa Sulphurata, Potassii Acetas, Potassii Bicarbonas, Potassii Citras and Potassii Tartras.

Foreign Pharmacopœias—Official in all, Austr, Belg, Ger, Hung, Jap and Russ (Kalium Carbonicum), Dan, Dutch, Norw and Swed (Carbonas Kalicus), Fr (Carbonate Neutre de Potassium), Ital (Carbonato di Potassio), Mex (Carbonato de Potasio Neutro), Port (Carbonato de Potassa), Span (Carbonato Potasico), Swiss has (Kalium Carbonicum Depuratum) and (Kalium Carbonicum Purum), US (Potassii Carbonas), Austri, Dutch, Ger, Jap, Russ and Swed, include a crude Carbonate, Ger, a 33½ p c Liquor, Swed, a solution, 20 p c

Tests—Potassium Carbonate, *BP*, when heated at a red heat loses from 15.0 to 17.0 p c, and leaves between 83.0 to 85.0 p c of anhydrous Potassium Carbonate. The *USP* states that when heated to 130° C (266° F) the salt loses all the Water which it may have retained or absorbed. It dissolves readily in Water, forming a clear solution which has a strong alkaline reaction towards red Litmus paper, which, when neutralised with Hydrochloric Acid, yields the tests distinctive of Potassium given under that heading, and which, on the addition of diluted acids, effervesces, evolving a colourless and odourless gas, which, when passed into Lime Water, yields a white precipitate soluble in a sufficient excess of the gas and redissolving with effervescence in diluted acids. It is officially required to contain 81.6 p c of pure Potassium Carbonate as determined volumetrically by direct titration with Volumetric Sulphuric Acid Solution as indicated below. The *USP* requires that it shall contain not less than 98 p c of pure Potassium Carbonate as volumetrically determined by direct titration, using Methyl Orange T.S. as an indicator of neutrality, the process is given below. The *PG* requires that it shall contain at least 95 p c of pure Potassium Carbonate as volumetrically determined by direct titration as given below.

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead and Zinc, Aluminium and Magnesium, Chlorides, Nitrates and Sulphates. The *BP* also includes Sodium, Cyanides, Sulphides and Thiosulphates. When acidified with Hydrochloric Acid the aqueous solution should yield no reaction with the modified Gutzzeit's test. It should yield no reaction with the Hydrogen Sulphide test described below. It should give no flocculent precipitate nor a turbidity when made slightly alkaline with Ammonia Solution and boiled, indicating the absence of Aluminium. The filtrate from this treatment should neither yield a marked turbidity nor a precipitate with Ammonium Oxalate Solution, indicating the absence of more than traces of Calcium. If this liquid be again filtered it should neither yield a pronounced turbidity nor a precipitate with Sodium Phosphate Solution, indicating the absence of more than traces of Magnesium. The aqueous solution, when acidified with Nitric Acid, should yield only a slight turbidity with Silver Nitrate Solution, indicating the absence of more than traces of Chlorides. It should answer the tests for freedom from Nitrate with Ferrous Sulphate and Sulphuric Acid given below, and also the test with Barium Nitrate Solution. When moistened with Hydrochloric Acid and inserted on a platinum loop into a non-luminous flame

it should communicate to the flame at most but a transient yellow coloration. On the addition of Sulphuric Acid in slight excess it should not evolve an odour of Hydrocyanic Acid, nor should the issuing gas cause a piece of filter paper moistened with Lead Acetate Solution to darken in colour when suspended in the gas, indicating the absence of Sulphides. Neither should the issuing gas possess an odour of Sulphur Dioxide nor bleach a piece of blue Litmus paper suspended therein, the liquid, after the addition of slight excess of Hydrochloric Acid, should remain clear, indicating the absence of Thiosulphate. The *U S P* gives a test for earthy impurities, requiring that no residue should remain when 1 gramme of the salt is dissolved in 20 c.c. of Water.

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by T.S. of Hydrogen Sulphide, either before or after acidification with Acetic Acid, *P G*, an aqueous solution (1-20) slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *U S P*.

Silver Nitrate—1 volume of a (1-20) aqueous solution should give with 10 volumes of Tenth-normal Volumetric Silver Nitrate Solution a yellowish-white precipitate which on gently warming does not become darker in colour, *P G*, a (1-20) aqueous solution saturated with diluted Nitric Acid should not be rendered more than opalescent after 2 minutes by T.S. of Silver Nitrate, *P G*.

Barium Nitrate—An aqueous solution (1-20) saturated with Acetic Acid should not be affected by T.S. of Barium Nitrate, *P G*.

Potassium Ferrocyanide—20 c.c. of an aqueous solution (1-20) over-neutralised with Hydrochloric Acid should not be rendered blue by 0.5 c.c. of T.S. of Potassium Ferrocyanide, *P G*.

Ferrous Sulphate and Ferric Chloride.—The 1-20 aqueous solution of the salt when mixed and gently warmed with a little Ferrous Sulphate T.S., and Ferric Chloride T.S. should not develop a blue colour on the addition of an excess of Hydrochloric Acid, *P G*, indicating the absence of Cyanide.

Ferrous Sulphate and Sulphuric Acid—2 c.c. of a solution of the salt in diluted Sulphuric Acid mixed with 2 c.c. of Sulphuric Acid and 1 c.c. of Ferrous Sulphate T.S. poured upon it as a layer should not give any coloured zone, *P G*, 8 c.c. of Ferrous Sulphate T.S. carefully poured upon 5 c.c. of a cold aqueous (1-20) solution of the salt mixed with 5 c.c. of Sulphuric Acid should not develop a brown colour at the junction of the two liquids, *U S P*.

Volumetric Determination—Not less than 11.9 c.c. of Volumetric Sulphuric Acid Solution should be necessary to neutralise 1 gramme of the salt, *B P*, 1 gramme of the salt should require for neutralisation not less than 13.7 c.c. of Normal Volumetric Solution of Hydrochloric Acid, *P G*, 1 gramme of the salt thoroughly dried at 130° C (266° F) dissolved in 50 c.c. of Water should require not less than 14.8 (14.28) of Normal Sulphuric Acid Volumetric Solution for neutralisation, using Methyl Orange T.S. as indicator, *U S P*.

POTASSII CHLORAS.

POTASSIUM CHLORATE

KClO_3 , eq 121.66

FR, CHLORATE DE POTASSIUM, GER, KALIUMCHLORAT, ITAL, CLORATO DI POTASSIO, SPAN, CLORATO POTASICO

Colourless, glistening, translucent, monoclinic prisms or plates, or as a white, odourless powder, possessing a cooling saline taste. It should be kept in well-stoppered bottles.

On account of the ready manner in which it evolves Oxygen, it should be handled with caution, great care being taken to avoid friction or any sudden percussion when mixing it with readily oxidisable or inflammable substances. When triturated with certain substances, *e g*, Sulphur, Sugar, Tannic Acid and Antimony Sulphide, it forms explosive mixtures. It has also been known to explode whilst being compressed into tablets.

Solubility—1 in 16 of cold Water, 1 in 2 of boiling Water, 1 in 1700 of Alcohol (90 p c), 1 in 152 of Alcohol (60 p c)

Medicinal Properties—A local stimulant. A strong solution, 1 or 2 in 40 of Water, is the best **wash** for the mouth when the gums are spongy, inflamed and irritable, and for ulcerative stomatitis, it relieves the tenderness and induces a firmness of the gums, it is also an excellent **gargle** in tonsillitis. The powder is applied to aphthæ in the mouth. Internally it is given to prevent the tendency to miscarriage, and to foetal death. In young people it should be used with great care and in small doses, if given at all.

Dose—5 to 15 grains = 0.32 to 1 gramme

10 grains 3 times daily for 6 months with no ill effects, in habitual death of the fetus in the later months of pregnancy.—*L* '02, ii 459

Hyperplasia of the foetal thyroid in cases where the mother had been given Potassium Chlorate.—*B M J* '03 i 657, 874

As a galactagogue, *T G* '93, 322, internally 7 drms taken by mistake caused death.—*L* '79, i 206

40 to 60 grains each of this salt and Carbolic Acid in 8 oz of Water will be found a pleasant and efficacious solution with which to brush the teeth and wash out the mouth and throat as a preliminary treatment in neurasthenia.—*B M J* '06, i 493

Incompatibles—Charcoal, Sulphur and Ferrous salts. Hydrochloric Acid causes the evolution of Chlorine, other mineral acids, of various chlorous smelling oxy-compounds, organic acids the same but much more slowly.

Official Preparation—Trochiscus Potassii Chloratis, used in the preparation of Potassii Permanganas.

Not Official—Gargarisma Potassii Chloratis, Gargarisme au Chlorate de Potassium, Mistura Potassii Chloratis, Pulvis Potassii Chloratis Compositus and Sodu Chloras.

Foreign Pharmacopœias—Official in Austr, Belg, Ger, Hung, Jap, Russ and Swiss (Kalium Chloricum), Dan, Dutch, Norw and Swed (Chloras Kalicus), Fr (Chlorate de Potassium), Ital (Clorato di Potassio), Mex (Clorato de Potasio), Port (Chlorato de Potassa), Span (Clorato Potasico), US.

Tests—Potassium Chlorate fuses when strongly heated, evolving a colourless and odourless gas, if the glowing end of a splinter of wood be inserted into the containing vessel it immediately ignites. When heated till no further gas is evolved it leaves a white residue, and if this residue be dissolved in Water a solution is yielded which, on the addition of Silver Nitrate Solution, affords a white precipitate, insoluble in Nitric Acid, readily soluble in Ammonia Solution and in Potassium Cyanide Solution. The aqueous solution of this residue also affords the tests distinctive of Potassium given under that heading. The *USP* states that it fuses at 334° C (633 2° F) and decomposes above 352° C (665 6° F), the whole of the Oxygen being evolved above 400° C (752° F). It also states that the amount of

residue left amounts to 60.8 p.c. of the pure Chlorate employed. The salt, when treated with Hydrochloric Acid and warmed, evolves a yellow gas possessing a strong chlorinous odour and stated by the *B.P.* to be a mixture of Chlorine and Chloric Oxide. It dissolves in Water, the resulting solution having a neutral reaction towards Litmus paper.

The more generally occurring impurities are Aluminium, Calcium, Copper, Iron, Lead, Magnesium and Sodium, Chlorides, Nitrates and Sulphates. The aqueous solution should neither yield a turbidity nor a flocculent precipitate when boiled with Ammonia Solution. An aqueous solution of the salt should yield neither a turbidity nor a precipitate with Ammonium Oxalate Solution. It should not be affected by the Hydrogen Sulphide or Ammonium Sulphide test described below. A standard has been suggested (*C.D.* '08, 1796) of 10 parts per million for Lead and 2 parts per million for Arsenic. If the liquid, to which Ammonium Oxalate Solution is added, be filtered it should yield neither a turbidity nor a precipitate on the addition of Sodium Phosphate Solution. A crystal of the salt when moistened with Hydrochloric Acid and inserted into a non-luminous flame on a loop of platinum wire should not afford a distinct or permanent yellow coloration to the flame. The aqueous solution should not afford more than a faint turbidity on the addition of Silver Nitrate or Barium Chloride Solution. It is officially required to yield no characteristic reaction with the tests for Nitrates, but in this instance the usual tests for Nitrates, unless carried out with considerable modification, are of no avail. In testing for Nitrates the *U.S.P.* employs Potassium Hydroxide T.S. and Aluminium wire. The *P.G.* employs a solution of Sodium Hydroxide with a mixture of Zinc filings and powdered Iron as described below.

Hydrogen Sulphide or Ammonium Sulphide.—An aqueous solution of the salt (1-20) should not be affected by T.S. of Hydrogen Sulphide, *P.G.*, should not become discoloured by T.S. of Ammonium Sulphide, *U.S.P.*

Ammonium Oxalate.—An aqueous solution of the salt (1-20) should not be affected by T.S. of Ammonium Oxalate, *P.G.*

Barium Nitrate.—An aqueous solution of the salt (1-20) should not be affected by T.S. of Barium Nitrate, *P.G.*

Silver Nitrate.—An aqueous solution of the salt (1-20) should not be affected by T.S. of Silver Nitrate, *P.G.*

Potassium Ferrocyanide.—20 c.c. of an aqueous solution of the salt (1-20) should not be rendered blue by T.S. of Potassium Ferrocyanide, *P.G.*

Sodium Hydroxide, Zinc and Iron Filings.—1 gramme of Potassium Chlorate warmed with 5 c.c. Sodium Hydroxide T.S. and a mixture of 0.5 gramme of Zinc filings and powdered Iron should not evolve Ammonia, *P.G.*

Potassium Hydroxide and Aluminium Wire.—If to 1 gramme of the salt contained in a test-tube of capacity, 5 c.c. of Water, 5 c.c. of Potassium Hydroxide T.S., and a small piece of Aluminium wire be added, and if in the upper portion of the test-tube a pledget of purified cotton be inserted, and over the mouth there be placed a piece of moistened red Litmus paper, then if the tube be heated upon a water-bath for 15 minutes, no blue coloration of the paper should be discernible, *U.S.P.*

Preparation

TROCHISCUS POTASSII CHLORATIS. POTASSIUM CHLORATE

LOZENGE

3 grains of Potassium Chlorate in each, with Rose Basis

Dose —1 to 6 lozenges

Potassium Chlorate is supplied in tablets or compressed discs, also combined with Borax and with Cocaine

Foreign Pharmacopœias — Official in Belg (Tabellæ), $1\frac{1}{2}$ grains, Dutch, $1\frac{1}{2}$ grains, Fr (Tablettes), $1\frac{1}{2}$ grains, Ital (Pastiglia), $1\frac{1}{2}$ grains, Jap (Pastilli), $1\frac{1}{2}$ grains, Mex (Pastillas), $1\frac{1}{2}$ grains, Port (Pastilhas), $1\frac{1}{2}$ grains, Span (Tabletas), $1\frac{1}{2}$ grains, Swiss (Pastilli), $1\frac{1}{2}$ grains, U S, about $2\frac{1}{2}$ grains in each lozenge

Not Official

GARGARISMA POTASSII CHLORATIS —Potassium Chlorate, 1 dr̄m , Glycerin, $\frac{1}{2}$ fl oz , Water, to 6 fl oz

Potassium Chlorate, 200 grains, Diluted Hydrochloric Acid, 100 minims, Water, to 20 fl oz —*St Thomas's*

This has been incorporated in the *B P C* as follows —Potassium Chlorate, 2, Diluted Hydrochloric Acid, 1, Water, to produce 100 —*B P C*

See also Gargarisma Chlori, p 371

GARGARISME AU CHLORATE DE POTASSIUM —Potassium Chlorate, 1, Distilled Water, 25, Syrup of Mulberries, 5, all by weight —*Fr*

MISTURA POTASSII CHLORATIS —Potassium Chlorate, 10 grains, Diluted Hydrochloric Acid, 5 minims, Distilled Water, to 1 fl oz —*St Thomas's*

This has been incorporated in the *B P C*

PULVIS POTASSII CHLORATIS COMPOSITUS — Potassium Chlorate, 1, Borax, 1, Sodium Bicarbonate, 1, White Sugar, 2, all in powder Mix A measured teaspoonful to be dissolved in half a tumbler (5 fl oz) of tepid Water, half the solution to be injected with a syringe along the floor of each nostril night and morning After use blow nose freely —*Central Throat*

SODII CHLORAS —Colourless, translucent crystals, or a white crystalline powder Soluble in about its own weight of Water, and in 5 times its weight of Glycerin

The same remarks with regard to caution in its use apply to this as to the Potassium salt

Official in Fr, Mex, Span and U S

POTASSII CITRAS

POTASSIUM CITRATE

$K_3C_6H_5O_7 \cdot H_2O$, eq 321.99

Fr, POTION GAZEUSE, GER, RIVER'SCHFR TRANK, ITAL, CITRATO DI POTASSIO, SPAN, CITRATO POTASICO

Translucent, prismatic crystals, or as a white, granular, deliquescent powder, possessing a cooling saline taste It should be preserved in well-stoppered bottles

The *B P* formula for this salt omits the molecule of Water of crystallisation The *U S P* gives the formula with 1 molecule of Water The salt is not official in the *P G*,

Solubility—10 in 6 of Water, and measures 11, 1 in 2 of Glycerin, 1 in 9 of Alcohol (60 p c), but if more of the salt is added the Alcohol separates from the watery solution

Medicinal Properties—Antacid, mild diaphoretic and diuretic, also alkalis the urine, and its free administration in acute nephritis is strongly advocated by Fothergill and others. Useful in gout and rheumatism, and in bronchitis with viscid, scanty expectoration. Given as a drink in scurvy.

Free administration combined with Colchicum in the treatment of gout—*L* '99, ii 1362

Milder cases of bacterial cystitis, pyelitis, and incontinence of urine in children do well with full doses of this combined with sedatives of the Belladonna group—*L* '08, i 79

In 15 grain doses in diabetic acetonaemia, because it is not neutralised by the gastric juice and becomes Bicarbonate in the blood—*P* '07, ii 120

Dose—10 to 40 grains = 0.65 to 2.6 grammes

Not Official—Mistura Potassii Citris Effervescens

Foreign P—Official in Port and U S. U S has also Potassii Citras in the others

Various solutions of Potassium Citrate occur as follows: Belg., Dutch, Hung. and Russ. (*Potio Riverii*), Fr. (*Potion Gazeuse*), Ger. (*Potio Riveri*, made with Sodium Carbonate and Citric Acid), Dan. and Norw. (*Julapium Salinum*), Port. (*Solutio de Citrato de Potassa*), Swiss (*Potio Effervescens*), U S. (*Liquor Potassae Citratis*)

Tests—Potassium Citrate when heated above 100° C (212° F) loses Water, and at 200° C (392° F), according to the *USP*, the Water of crystallisation is completely lost, the loss amounting to 5.55 p c. When heated to a still higher temperature it chars, and when ignited to a dull red heat leaves a mixture of Potassium Carbonate and Carbon. It dissolves very readily in Water, forming a solution which has an alkaline reaction towards red Litmus paper, but is neutral to Phenolphthalein Solution. The aqueous solution, when acidified with Hydrochloric Acid, yields the tests of Potassium given under that heading. It yields when boiled with an excess of Calcium Chloride Solution a white precipitate insoluble in Potassium Hydroxide Solution, but soluble in Ammonium Chloride Solution and in solutions of alkali Citrates; it yields with Silver Nitrate Solution a white precipitate soluble in Ammonia Solution, but in contradistinction to the precipitate yielded by Tartrates does not yield a mirror when the ammoniacal solution is warmed. It is officially required to contain at least 98.3 p c of Potassium Citrate of the pharminacopoeial formula, as volumetrically determined by titrating the solution of the residue left on ignition with Volumetric Sulphuric Acid Solution as shown below. It has been found that the volumetric requirements do not correspond to the formula given, and should be altered. The *USP* requires that it should contain not less than 99 p c of pure Potassium Citrate as volumetrically determined by titrating the solution of the residue left on ignition with Semi-normal Volumetric Hydrochloric Acid Solution, using Methyl Orange T S as an indicator of neutrality. The method is given in small type below.

The more generally occurring impurities are Calcium, Iron, Lead, Magnesium and Sodium, Carbonates and Tartrates, Chlorides and Sulphates. The aqueous solution should afford no reaction with the Hydrogen Sulphide test described below. A standard has been suggested (*C D '08*, 1796) of 5 parts per million for Lead and 1 part per million for Arsenic. It should afford neither a turbidity nor a precipitate with Ammonium Oxalate Solution, and the filtrate from the Ammonium Oxalate should afford neither a turbidity nor a precipitate with Sodium Phosphate Solution. A crystal of the salt when moistened with Hydrochloric Acid and brought into the non-luminous flame on a loop of platinum wire should at the most afford only a transient yellow coloration to the flame. The addition of diluted Hydrochloric Acid Solution should not cause effervescence in the concentrated aqueous solution. It should yield no reaction with the Acetic Acid test described below. The aqueous solution when acidified with diluted Nitric Acid should yield at the most a faint turbidity on the addition of Silver Nitrate or Barium Chloride Solution.

Hydrogen Sulphide—An aqueous solution of the salt (1.20) slightly acidulated with Acetic Acid should not respond to the time limit test for heavy metals, *U S P*.

Acetic Acid—1 gramme of the salt dissolved in 1 c.c. of Water should not deposit any precipitate on the addition of 1 c.c. of Acetic Acid, indicating absence of Tartrate, *U S P*.

Volumetric Determination—The filtered aqueous solution obtained by dissolving and filtering the residue left on igniting 1 gramme of dry salt at a red heat, should require for neutralisation not less than 9.7 c.c. of Volumetric Solution of Sulphuric Acid, *B P*. If 1 gramme of the salt be treated as described under Potassium Acetate it should require for neutralisation not less than 18.4 c.c. of Semi-normal Volumetric Hydrochloric Acid Solution, using Methyl Orange T.S. as indicator, *U S P*.

Not Official

MISTURA POTASSII CITRAS EFFERVESCENS—Potassium Bicarbonate, 20 grains, Water, to 1 fl. oz. (Alkaline Solution). Citric Acid, 15 grains, Water, to $\frac{1}{2}$ fl. oz. (Acid Solution). Mix the two solutions and drink during effervescence.—*St Thomas's*

This has been incorporated in the *B P C*

Not Official

POTASSII CYANIDUM

KCN, eq. 64.68

FR, CYANURE DE POTASSIUM, GER, KALIUMCYANID, ITAL, CIANURO DI POTASSIO, SPAN, CIANURO POTASSICO

White, opaque, deliquescent masses, or as a white, granular, deliquescent powder, having the odour of Hydrocyanic Acid. The pure salt can be obtained in white cubical crystals. It is intensely poisonous.

It should be kept in well stoppered bottles.

The commercial salt containing at least 90 p.c. of Potassium Cyanide is official in the Appendix to the *B P*.

Solubility—1 in $2\frac{1}{2}$ of Water, almost entirely 1 in 100 of Alcohol (90 p.c.)

Ordinary fused Cyanide only contains about 40 p.c. of real Cyanide, but there is no difficulty in obtaining it from 95 to 99 p.c.

It is useful to remove the black stains on the skin caused by Silver Nitrate. Entomologists use it with gypsum to make poison bottles for killing insects without injuring their delicate structure, for this purpose dissolve 1 of the Cyanide in $1\frac{1}{2}$ of Water, and add 2 of Plaster of Paris. This mixture stirred and poured whilst liquid into a wide-mouthed bottle, forms a hard floor, which is constantly giving off vapour.

Foreign Pharmacopœias—Official in Fr, Mex, Port, Span and U S Not in the others

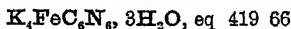
Tests—Potassium Cyanide fuses at a low red heat, it dissolves readily in Water, forming a clear solution which has a reaction towards red Litmus paper. When treated with a slight excess of diluted Hydrochloric Acid it evolves a highly poisonous and characteristic odour of Hydrocyanic Acid, the resulting solution affords the tests characteristic of Potassium given under that heading. The aqueous solution yields with Silver Nitrate T.S. a white, curdy precipitate soluble in excess of Potassium Cyanide Solution, in Solution of Ammonia and in concentrated Nitric Acid. When shaken with a few drops of a mixture of Ferrous Sulphate T.S. and Ferric Chloride T.S. it yields, on the addition of a slight excess of Hydrochloric Acid, a blue precipitate. No method of determination is given in the B.P. The U.S.P. requires that it shall yield not less than 95 p.c. of pure Potassium Cyanide as volumetrically determined by dissolving 1 gramme of the salt in sufficient Water to measure 100 c.c., mixing 64.7 c.c. of this solution with 5 c.c. of Ammonia and 3 drops of 20 p.c. w/v Potassium Iodide Solution and titrating the mixture with Tenth-normal Volumetric Silver Nitrate Solution, not less than 47.5 c.c. should be required to produce a permanent precipitate. 1 c.c. of Tenth-normal Volumetric Silver Nitrate Solution is equivalent to 2 p.c. of Cyanide.

The more generally occurring Ferrocyanide and Sulphocyanide. The addition of Hydrochloric Acid in slight excess to a 5 p.c. solution of the salt should not yield more than a slight effervescence, and the addition of 1 drop of Ferric Chloride T.S. to this liquid should neither produce a blue nor a red coloration.

Not Official

POTASSII FERROCYANIDUM

Syn—YELLOW PRUSSIAN OF POTASH



Large translucent, lemon-yellow, soft, table-shaped crystals or groups of crystals, efflorescent in dry air.

It is officially described as a yellow crystalline salt prepared by fusing together Potassium Carbonate, nitrogenous organic matter and Iron.

It should be kept in well-stoppered glass bottles of a dark amber tint and exposed as little as possible to the air.

Solubility—1 in 4 of Water, insoluble in Alcohol (90 p.c.)

Foreign Pharmacopœias—Official in Mex, Port, Span and U S Not in the others

Tests—Potassium Ferrocyanide loses its Water of crystallisation when heated, and is converted into an anhydrous salt at a temperature of 100°C (212°F), it dissolves readily in Water, forming a clear solution which is neutral in reaction towards Litmus paper. A concentrated solution yields with Sodium Hydroxide a white crystalline precipitate. Lead yields a white precipitate. It yields a reddish-brown precipitate with Silver Nitrate. It produces a dark blue precipitate. It may be distinguished from Ferrocyanide. A concentrated solution of the salt should show no effervescence on the addition of diluted Sulphuric Acid. The reaction of Silver Nitrate Solution to an aqueous solution acidified with Nitric Acid should yield a white precipitate free from a red tint.

Not Official

POTASSII HYPOPHOSPHIS

POTASSIUM HYPOPHOSPHITE

KPH O, eq 103.39

White, opaque crystals, or in crystalline masses, or as a white, granular deliquescent powder, possessing a pungent and saline taste.

It should be kept in well closed glass bottles in a cool atmosphere and protected as far as possible from contact with the air, as it is very deliquescent. It should be handled with great caution, as it is readily oxidized, and when brought into contact with substances which readily part with Oxygen the temperature rises so rapidly that an explosion often results.

Solubility—10 in 6 of Water, 1 in 7 of Alcohol (90 p c), 3 in 1 of boiling Water, 1 in 375 of Ether, insoluble in Chloroform.

Tests—Potassium Hypophosphite when heated in a dry tube loses moisture, and when more strongly heated evolves spontaneously inflammable Hydrogen Phosphide gas which burns with a bright yellow flame. It dissolves readily in Water, yielding a clear solution which is neutral towards Litmus paper, or only faintly alkaline towards red Litmus paper. It yields the tests distinctive of Potassium given under that heading. The diluted aqueous solution slightly acidulated with diluted Sulphuric Acid yields, on the addition of Silver Nitrate Solution, a white precipitate rapidly becoming brown or black, owing to its reduction to metallic Silver. The aqueous solution when gently warmed with Copper Sulphate Solution yields a reddish brown precipitate. The aqueous solution of the salt, when acidulated with Hydrochloric Acid and added drop by drop to an excess of Mercuric Chloride Solution, yields a white precipitate of Mercurous Chloride, and on the further addition of an excess of the solution of the salt the white precipitate becomes grey, owing to its reduction to metallic Mercury. Its aqueous solution readily reduces Potassium Permanganate, the purple colour of a Permanganate Solution being rapidly discharged. It is required by the *U S P* to contain not less than 98 p c of pure Potassium Hypophosphite, but no method of determination is given. The percentage of absolute Potassium Hypophosphite present may be readily ascertained by the method given under Calcium Hypophosphis.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Zinc, Calcium, Chlorides and Sulphates, Phosphates and Phosphites. Arsenic, if present, may be detected by the modified Gutzet's test, treating 5 c c of a 10 p c aqueous solution of the salt with 3 c c of Nitric Acid diluted with about 10 c c of water, evaporating the mixture to dryness on the water-bath, and performing the test on the residue. Copper, Lead, Iron and Zinc, if present, may be detected by Hydrogen Sulphide, either in a solution rendered faintly acid with Hydrochloric Acid, or in a solution made alkaline with Ammonia Solution. The aqueous solution of the salt should not afford a distinct turbidity with Ammonium Oxalate Solution after the addition of a little Ammonium Chloride Solution, indicating the absence of Calcium. If the mixture be allowed to stand for some time and filtered, it should yield little or no turbidity with Sodium Phosphate Solution, indicating the absence of Magnesium. When acidified with diluted Nitric Acid the aqueous solution should not afford a pronounced turbidity or precipitate with either Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. It should not yield a very pronounced turbidity or precipitate on the addition of Lead Acetate Solution, indicating a limit of Phosphates and Phosphites. Remarks on the Lead Acetate test will be found under Calcium Hypophosphis and under Sodium Hypophosphis. A 5 p c aqueous solution of the salt should not effervesce on the addition of an acid, indicating the absence of Carbonate.

POTASSII IODIDUM.

POTASSIUM IODIDE

KI, eq 164 73FR, IODURE DE POTASSIUM, GER, KALIUMJODID, ITAL, JODURO DI POTASSIO
SPAN, YODURO POTASICO

Colourless translucent, or white opaque, cubical crystals, permanent in dry air. It has a characteristic saline, subsequently somewhat bitter, metallic taste, and a faintly alkaline reaction.

It should be preserved from air and light in well-stoppered bottles and kept in a cool place.

Solubility—4 in 3 of Water, and measures 4, 1 in 10 of Alcohol (90 p c), 1 in 3 of Glycerin.

Medicinal Properties—Alterative, diuretic, expectorant. It is useful in cases where Iodine is indicated, and being less irritating is much preferred for internal administration. Useful especially in secondary and in tertiary syphilis and in all diseases associated with syphilis, such for example as locomotor ataxy. For secondary symptoms 60 grains in solution may be given in the 24 hours. It reduces chronic inflammatory swelling, effusions and glandular enlargement, and is useful in goitre and obesity, also in chronic Bright's disease, bronchitis and bronchitic asthma, aortic disease, endocarditis, internal aneurism, angina pectoris (the pain of which it also relieves), and in established arterio-sclerosis, chronic rheumatism and gout, lumbago, sciatica, psoriasis and, in large doses (*see below*), actinomycosis. May be given with Quinine dissolved by Sulphuric or Phosphoric Acid, but not with Nitro-hydrochloric Acid, as the eliminated Chlorine decomposes it and makes an unsightly mixture. Combined with Nux Vomica the system bears it better. It is useful in the elimination of Lead in cases of chronic lead poisoning, also in treating chronic mercurial poisoning. *See also under 'Iodum'*.

(*B M J* '04, ii 1068) its efficacious use in rheumatism which was frequently the cause of ovarian pain. Its use in actinomycosis yields little improvement under a dose of less than 20 grains thrice daily (*L* '04, ii 1225), and in some instances 1-dr. doses thrice daily have been recommended, but these should be given with large quantities of Water. Sodium Iodide is preferable on account of depression caused by the Potassium salt. It has been recommended (*B M J* '04, ii 1206) in the internal treatment of non-suppurative middle-ear disease. In large doses (*B M J* '04, ii 1209), to check the ossifying process in the early stages of otosclerosis.

Should always be given in solution well diluted, and if possible never on an empty stomach. Milk is the best diluent. The drug should never be given in phthisis.—*B M J E* '05, ii 15.

Where rigid arteries were a cause of insomnia the use of massage and the administration of this salt were of especial value.—*B M J* '05, ii 249.

Indicated in the treatment for ordinary cases of Yaws (frambæsia), in large doses.—*L* '07, ii 1459.

In cretinism.—*L* '93, ii 1545.

Sodium or Potassium Iodide when given to man by the stomach in ordinary doses has no depressing effect on the action of the heart, or on the blood pressure in the arteries.—*B M J* '01, ii 1524.

The opinion is expressed that Iodides are of no value in the treatment of aneurism, and that they are even hurtful in arterio-sclerosis, but the great bulk

of medical opinion and experience is directly contrary to this—*B M J* '01, ii 1522

Productive of good results when uterine sclerosis is established, but absolutely contra indicated in the pre sclerosis stage—*B M J E* '07, i 83

Its regular use in small doses, with the occasional exhibition of *Strophanthu*, often gives considerable relief in cardio arterial disease—*B M J* '01, ii 1057

60 to 80 grains 3 times daily, successful in thoracic aneurisms—*L* '03, ii 528

Good results from its prolonged administration combined with one of the guaiacol preparations in acute rheumatoid arthritis—*B M J* '01, ii 1039

Cases where congenital goitre followed the administration of Potassium Iodide to the mother during pregnancy—*L* '07, i 1714

Case of Iodine rash after administration of 3 doses of 10 grains each—*L* '04 i 421

Dose—5 to 20 grains = 0.32 to 1.3 grammes, this dose is often greatly exceeded, especially in syphilis of the nervous system

Prescribing Notes—*Best given with Tincture of Orange and Spirit of Chloroform, in Water, or with Tincture of Cinchona. It is also given with Fowler's Solution to prevent the rash sometimes produced. Tablets cause gastric pain. Solutions of Ferric salts, when acid, set free Iodine from Potassium Iodide.*

It is better borne when given with Potassium Acetate, or when administered alternately with Ferrous Iodide—*L* '88, i 1019

Incompatibles—*Spiritus Ætheris Nitrosi, Bismuthi Subnitras*

Official Preparations—*Linimentum Potassii Iodidi cum Sapone* and *Unguentum Potassii Iodidi*, contained in *Liquor Iodi Fortis, Tinctura Iodi* and *Unguentum Iodi*. Used in the preparation of *Hydrargyri Iodidum Rubrum* and *Plumbi Iodidum*

Not Official—*Linimentum Potassii Iodidi, Linimentum Potassii Iodidi cum Sapone (B P '67), Mistura Potassii Iodidi, Mistura Potassii Iodidi Alkalina, Mistura Potassii Iodidi et Stramonii, Pilulæ Kalii Iodati, Pomada de Yoduro Potassico con Extracto de Cicuta*

Foreign Pharmacopœias—Official in Austr, Belg, Ger, Hung, Jap, Russ and Swiss, (*Kalium Iodatum*), Dan, Dutch, Norw and Swed, (*Iodetum Kalicum*), Fr, (*Iodure de Potassium*), Ital, (*Ioduro di Potassio*), Mex, (*Yoduro de Potasio*), Port, (*Iodeto de Potassio*), Span, (*Yoduro Potassico*), U S

Tests—Potassium Iodide when heated decrepitates, and when strongly heated fuses. It dissolves readily in Water, forming a clear solution which possesses a faintly alkaline reaction towards red Litmus paper, and which yields the tests distinctive of Potassium given under that substance. It produces with Silver Nitrate Solution a yellow, curdy precipitate insoluble in Nitric Acid, insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution. With Mercuric Chloride Solution it yields a brilliant scarlet precipitate slightly soluble in excess of the reagent and readily soluble in the Iodide Solution. With Lead Acetate it yields a yellow crystalline precipitate soluble in diluted Nitric Acid and also in boiling Water, it is deposited, as the solution cools, in brilliant golden crystalline scales. When cautiously mixed with a little Chlorine Water or Bromine Solution it yields a brown or reddish-brown coloration, and on the addition of Starch Mucilage an intense blue colour, if the brown coloured solution be shaken with Carbon Bisulphide, the Bisulphide Solution assumes a deep violet colour. It is officially required to contain not less than 98.0 p.c, nor more than 101.9 p.c. of pure Potassium Iodide as volu-

metrically determined by direct titration of the salt with Volumetric Silver Nitrate Solution. The *USP* requires that the salt should contain at least 99.0 p.c. of pure Potassium Iodide as volumetrically determined by the direct titration of a solution of a well-dried salt in Water as indicated below. The *PG* does not give a requisite percentage of pure Potassium Iodide nor a method of determination.

The more generally occurring impurities are Arsenic, Aluminium, Calcium, Copper, Iron, Lead, Magnesium and Sodium, Bromides and Bromides, Carbonates, Chlorides, Cyanides, Iodates, Nitrates or Sulphates. The *BP* requires that it shall yield only the slightest reactions with the tests for Bromides, Carbonates, Chlorides or Sulphates, and no characteristic reaction for the remaining substances. The *USP* includes a test for limit of alkali, 'less soluble salts,' Barium and Thiosulphates. Arsenic, Copper, Lead and Iron may be detected, if present, by the Hydrogen Sulphide test described below, either before or after the addition of Ammonia. Neither a turbidity nor a flocculent precipitate should be produced when an aqueous solution of the salt is boiled with Ammonia Water. Ammonium Oxalate Solution should produce neither a turbidity nor a precipitate, nor should the addition of Sodium Phosphate Solution to the filtrate from the Ammonium Oxalate Solution cause any alteration. A crystal of the salt when moistened with Hydrochloric Acid and introduced into a non-luminous flame on a loop of platinum wire should not impart a persistent yellow coloration to the flame. When boiled with Liquor Potassæ an aqueous solution should not evolve an odour of Ammonia, nor should the issuing gas produce an alkaline reaction towards red Litmus paper. The aqueous solution should not afford an immediate yellow coloration when mixed with a little diluted Sulphuric Acid, when mixed with Chlorine Water the aqueous solution should afford when shaken with Carbon Bisulphide a distinct violet and not a brown coloration. A solution of the salt should not yield a pink coloration on the addition of Phenolphthalein Solution. Chlorides, if present, may be detected by shaking. Cyanides, if present, may be detected by the addition of an excess of Silver Nitrate Solution. With Ammonia Solution, the filtrate should yield no decided turbidity nor a precipitate on acidification with diluted Nitric Acid.

The Ferrous Sulphate and Potassium Hydroxide test described below may be employed for the detection of Cyanides. In testing for Nitrates the *USP* employs the Potassium Hydroxide and Aluminium wire test described in small type below. The *PG* the Sodium Hydroxide Solution, Zinc filings and powdered Iron test. Chlorides, Bromides and Thiosulphates, if present, may be detected by the test with Volumetric Silver Nitrate Solution mentioned in small type. Barium, if present, may be detected by the Potassium Sulphate test described below.

A method has been described (*PJ* '00 ii 58) by which the Iodine only, in a mixture of Chloride, Bromide and Iodide of Potassium, may be determined. The process depends upon the liberation of Iodine alone from the mixture by the interaction of 5 p.c. Potassium Bichromate Solution and 10 p.c. Sulphuric Acid Solution. A weighed

quantity of 0.5 of a gramme of a mixture of the salts is dissolved in 20 cc of Water, and 10 cc of a 5 pc Potassium Bichromate Solution is introduced, together with 10 cc of a 10 pc Sulphuric Acid Solution, the mixture is allowed to stand for a few minutes, shaking vigorously with 60 cc of Toluol. After separation the lower stratum is removed, the Toluol washed by agitation with small quantities of Water, adding the washings to the first portion separated. The mixed washings are again extracted with Toluol, which, if it be coloured violet, is added to that already separated. The mixed Toluol Solutions are then shaken with 35 cc of Tenth-normal Volumetric Sodium Thiosulphate Solution, the Thiosulphate Solution removed, the Toluol washed, the washings mixed with the Thiosulphate Solution, and the excess of Tenth-normal Volumetric Sodium Thiosulphate Solution titrated with Tenth-normal Volumetric Iodine Solution, from the amount of Tenth-normal Volumetric Sodium Thiosulphate Solution absorbed, the amount of Potassium Iodide may be readily calculated.

Litmus—Crushed Potassium Iodide brought in contact with moistened red Litmus paper should not immediately colour it violet blue, *P G*. The aqueous solution is neutral or has a slightly alkaline reaction upon Litmus paper *U S P*.

Phenolphthalein—A solution of 1 gramme of the salt in 10 cc of Water with 0.1 cc of Tenth-normal Volumetric Sulphuric Acid Solution added should yield no coloration on the subsequent addition of a drop of Phenolphthalein TS, even after heating, *U S P*.

Sulphuric Acid—A solution of 0.5 gramme of the salt in 10 cc of previously boiled and cooled Distilled Water, with the addition of 2 drops of diluted Sulphuric Acid (free from Sulphurous and Nitrous Acids) should show no distinct yellow colour within half a minute, *U S P*. The boiled and cooled (1-20) aqueous solution should not be immediately coloured on the addition of Starch Solution and diluted Sulphuric Acid.

Hydrogen Sulphide—The aqueous solution (1-20) should not be affected by TS of Hydrogen Sulphide, *P G*, slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *U S P*.

Barium Nitrate—An aqueous solution (1-20) should not be affected by TS of Barium Nitrate, *P G*.

Potassium Sulphate—10 cc of an aqueous solution (1-20) acidulated with Hydrochloric Acid should not be rendered turbid by 1 cc of TS of Potassium Sulphate, *U S P*.

Potassium Ferrocyanide—20 cc of an aqueous solution (1-20) acidulated with a few drops of Hydrochloric Acid should not be rendered blue by 0.5 cc of TS of Potassium Ferrocyanide, *P G*.

Ferrous Sulphate and Potassium or Sodium Hydroxide—If an aqueous solution (1-20) be gently warmed with a crystal of Ferrous Sulphate and 1 drop of Ferric Chloride TS after the addition of Sodium Hydroxide TS, it should not be coloured blue when saturated with Hydrochloric Acid, *P G*. The *U S P* directs that 5 cc of the aqueous solution be gently heated with 1 drop of Ferrous Sulphate TS and 0.5 cc of TS of Potassium Hydroxide, no blue colour should appear when the mixture is acidulated with Hydrochloric Acid.

Aluminium Wire and Potassium Hydroxide, or Zinc, Iron and Sodium Hydroxide—If 1 gramme of the salt be treated as described under Potassium Chlorate with Potassium Hydroxide TS and Aluminium wire the Litmus paper should show no blue coloration, *U S P*. 1 gramme of the salt

warmed with 5 c c of Sodium Hydroxide T S and a mixture of 0.5 gramme of Zinc filings and powdered Iron should not evolve Ammonia, *P G*

Volumetric Silver Nitrate and Ammonia Solution—If 0.2 gramme of the salt dissolved in 2 c c of Ammonia T S (10 p c *U S P*) be mixed with 13 c c of Tenth-normal Silver Nitrate Volumetric Solution the mixture agitated and filtered, the filtrate after acidulation with Nitric Acid should not become opaque (more than slightly turbid, *U S P*) nor should there be any darkening in colour within 10 minutes, *P G* and *U S P*

Volumetric Determination—Not less than 59.5 and not more than 61.9 c c of Silver Nitrate Volumetric Solution should be necessary to completely precipitate 1 gramme of the salt, *B P*, 0.5 gramme of the well-dried salt dissolved in 10 c c of Distilled Water and 3 drops of Potassium Chromate T S added should require not more than 30.8 c c and not less than 30 c c of Tenth normal Silver Nitrate Volumetric Solution to produce a permanent red colour, *U S P*

Preparations

LINIMENTUM POTASSII IODIDI CUM SAPONE. LINIMENT OF POTASSIUM IODIDE WITH SOAP

Curd Soap, recently prepared and in shavings, 2 oz, Potassium Iodide, 1½ oz, Glycerin, 1 fl oz, Oil of Lemon, 1 fl dr, Distilled Water, 10 fl oz. Dissolve the Soap and the Glycerin in the Water by the heat of a water-bath and pour the solution on to the powdered Potassium Iodide in a mortar, rub them together until the mixture is cold, after half an hour thoroughly mix in the Oil of Lemon. There will be a variable loss of Water in dissolving the Soap, and it should be made up to a weight when taken off the water-bath.

When first prepared it is very bulky, but after it has been made some time it occupies a much smaller space, and this is apt to cause trouble with patients. The difference is due to the quantity of air incorporated in it by the trituration, and is so great that it would be quite possible at different times for the same weight of Liniment to fill a 1 oz pot and a 4 oz pot.

The advantages of this Liniment are that it does not stain nor does it irritate when rubbed on the skin, it is employed in enlargement of the joints, and in indurated glands, especially the cervical glands.

Foreign Pharmacopœias—Official in Swiss (*Opodeldoc Iodatum*), Lard or Butter, 50, Alcohol (95 p c), 25, Solution of Caustic Soda, 25, saponify and dissolve in Alcohol, 825, Sodium Iodide, 50, Water, 25, Oil of Lemon, 10. Swiss has also *Opodeldoc Iodatum Liquidum*. Not in the others.

Linimentum Saponato-Camphoratum cum Kalio Iodato—Stearic Soap, 75, Vegetable Oil, 75, Spirits of Wine, 600, Water, 98, Glycerin, 50, Potassium Iodide, 100, Lavender Oil, 2—Aust.

UNGUENTUM POTASSII IODIDI POTASSIUM IODIDE OINTMENT

Potassium Iodide, 50, Potassium Carbonate, 3, Distilled Water (by weight), 47, Benzoated Lard, 400. Add the solution to the Lard in a slightly warmed mortar. (1 in 10)

Foreign Pharmacopœias—Official in Dutch, Fr, Ger, Hung, Ital, Norw, Port, Russ, Span, Swiss and U S, 1 in 10, Austr, Ger, Jap and Russ, with Sodium Thiosulphate. Mex, (*Pomada de Yoduro de Potasio*) Potassium Iodide, 8, Benzoated Lard, 60, Water, q s. Not in Belg or Dan.

Not Official.

LINIMENTUM POTASSII IODIDI.—Soft Soap, 2 oz, Potassium Iodide, 1½ oz, Glycerin, 1 fl oz, Oil of Lemon, 1 fl dr, Alcohol (60 p c), 10 fl oz—*Y B P* '04, 510.

These quantities correspond to those of the official Liniment, but Soft Soap and Alcohol (60 p c) are used in place of Curd Soap and Distilled Water.

It contains twice as much Soap as Linimentum Saponis
Soft Soap, 13 50, Potassium Iodide, 10, Glycerin, 7, Oil of Lemon, 1,
Alcohol (60 p c), *q s* to produce 100 —*B P C*

LINIMENTUM POTASSII IODIDI C SAPONE (*B P* 1867) Hard Soap,
1½ oz, Potassium Iodide, 1½ oz, Glycerin, 1 fl oz, Oil of Lemon, 1 fl drm
Water, 10 fl oz Put the Glycerin, Iodide, and 9 fl oz of Water in a clean
20 oz wide mouthed bottle, then dissolve the Soap (in shavings) in the 7 fl oz of
Water in a jar by the heat of a water bath, strain the solution whilst hot
through muslin into the bottle containing the Iodide, etc, allow to stand for
2 or 3 minutes until the bottom of the Soap Solution is a little opaque,
then mix by agitation, lastly add the Oil of Lemon, shaking briskly, and, after
agitating at intervals for 2 hours or more, a liniment in the form of a soft
white jelly will result, and remain so, if it should not, a small addition of Water
will generally perfect it

This formula is that of *B P* '67 but the manipulation has been modified,
when made properly it gives satisfaction

MISTURA POTASSII IODIDI —Potassium Iodide, 10 grains, Potassium
Bicarbonate, 5 grains, Pimento Water to 1 fl oz —*Brompton*

MISTURA POTASSII IODIDI ALKALINA —Potassium Bicarbonate,
15 grains, Ammonium Carbonate, 3 grains, Potassium Iodide, 3 grains,
Camphor Water to 1 fl oz —*Brompton* Potassium Iodide, 3 grains, Potassium
Bicarbonate, 10 grains, Ammonium Carbonate, 3 grains, Camphor Water, to 1
fl oz —*St Thomas's*

This has been incorporated in the *B P C*

MISTURA POTASSII IODIDI ET STRAMONII —Extract of Stramo-
num, ¼ grain, Extract of Liguonica, 2 grains, Potassium Iodide, 3 grains,
Chloric Ether, 5 minims, Water to 1 fl oz —*Brompton* Potassium Iodide,
3 grains, Tincture of Stramonium, 5 minims, Liquid Extract of Liguonica,
10 minims, Emulsion of Chloroform, 10 minims, Water, to 1 fl oz —*St Thomas's*

This has been incorporated in the *B P C*

PILULÆ KALII IODATI —Potassium Iodide, 20, Powdered Starch, 5,
Simple Syrup, *q s* —*Belg*

POMADA DE YODURO POTASICO CON EXTRACTO DE CICUTA
—Potassium Iodide Ointment, 80, Extract of Conium, 10, Distilled Water, *q s*
—*Span*

POTASSII NITRAS.

POTASSIUM NITRATE

B P Syn —NITRI, SALTPETRI

NO Syn —AZOTATE DI POTASSIO

KNO₃, eq 100 41

FR, AZOTATE DI POTASSIUM GER, KALIUMNITRAT, ITAL, NITRATO DI
POTASSIO, SPAN, NITRATO POTASICO

Colourless, transparent, rhombic prisms, or a white odourless,
crystalline powder, having a cooling saline taste It is obtained by
purifying Crude Nitre, or from Sodium Nitrate

It should be kept in well-stoppered glass bottles, as it has a slight
tendency to deliquesce in moist air

Solubility —1 in 4 of cold Water, 2½ in 1 of boiling Water,
sparingly in Alcohol (90 p c)

Medicinal Properties.—Sometimes given as a diuretic and
diaphoretic, but the Acetate and Citrate are much to be preferred

Useful as a gargle in relaxed sore throat Potassium Nitrate 5 grains, Potassium Bicarbonate 20 grains, taken, during effervescence, with Citric Acid 15 grains, in a small tumbler of cold Water, is a draught in febrile conditions A common ingredient of asthma inhalation powders Charta Nitrata is used in spasmodic asthma

15 grains of Potassium Nitrate with $\frac{1}{2}$ grain Sodium Nitrite useful for lessening high arterial tension and arresting epistaxis, 15 grains of Potassium Nitrate along with Potassium Bicarbonate and $\frac{1}{2}$ grain of Sodium Nitrite, useful for the same purpose in gouty subjects—*L* '02, 11 331, *B M J* '02, 11 504

Used for its slight antiseptic effect to assist in preserving canned meat, and to give it a red colour Amount should be restricted to 2 grains per pound—*Roy Army Med Corps Jour* '08, 1 124

Dose—5 to 20 grains = 0.32 to 1.3 grammes

Official Preparations—Contained in Argenti Nitras Induratus and Argenti Nitras Mitigatus Used in the preparation of Acidum Nitricum

Not Official—Mistura Salina, Mistura Salina Anodyna, Sal Prunella, Charta Nitrata, Charta Nitrata et Chlorata

Foreign Pharmacopœias—Official in all, Austr, Belg, Ger, Hung, Jap., Russ and Swiss (Kalicum Nitricum), Dan, Dutch, Norw and Swed (Nitras Kalicus), Fr (Azotate de Potassium), Ital (Nitrato di Potassio), Mex (Nitrato de Potasio), Port (Azotato de Potassa), Span (Nitrato Potasico), US (Potassii Nitras)

Tests—Potassium Nitrate when heated fuses The *USP* states at a temperature of 353° C (667.4° F), when still more strongly heated it is decomposed, giving off Oxygen and leaving a residue of Potassium Nitrate, Nitrite and Oxide It dissolves readily in Water, forming a clear solution which is neutral in reaction towards Litmus paper, and which yields the tests given under Potassium An aqueous solution when cautiously mixed with Sulphuric Acid, keeping the mixture cool, affords when a solution of Ferrous Sulphate is carefully poured on to the surface of the mixture a dark brown ring at the junction of the two fluids 1 or 2 drops of a Diphenylamine Solution (prepared by dissolving 0.1 of a gramme of Diphenylamine in 50 c.c. of Diluted Sulphuric Acid) when mixed with an aqueous solution affords a deep blue colour at the point of contact of the two liquids when Sulphuric Acid is carefully poured into the liquid, so as to form a separate layer An aqueous solution acidified with Sulphuric Acid immediately discharges the colour of indigo solution, when warmed with Sulphuric Acid and Copper foil an evolution of nitrous fumes ensues The *USP* states that the salt shall contain not less than 99 p.c. of pure Potassium Nitrate, but gives no method of determination Neither the *BP* nor the *PG* states a requisite percentage nor includes a method of determination

The more generally occurring impurities are Aluminium, Ammonium, Calcium, Copper, Iron, Lead, Magnesium, Sodium, Zinc, Chlorides, Iodides and Sulphates The *USP* includes tests for Chlorate and Perchlorate The aqueous solution of the salt should afford neither a turbidity nor a precipitate when boiled with Ammonia Solution It should yield no reaction for Arsenic, Copper, Iron, Lead or Zinc when examined by the Hydrogen Sulphide test described in

small type below It should not yield a reaction for Calcium or Magnesium when examined by the Ammonium Oxalate and Sodium Phosphate tests described below, nor for Chloride, Sulphate, Iodide, Chloride or Perchlorate when examined by the tests under Silver Nitrate, Barium Nitrate, Chlorine and Chloroform, and Sulphuric Acid given in the small type below The *P G* includes a separate test for Iron with Potassium Ferrocyanide Solution

Flame—Heated on Platinum wire it colours the flame violet, only a transitory yellow colour appearing, *P G*

Hydrogen Sulphide—The aqueous solution (1-20) should not be affected by T S of Hydrogen Sulphide, either before or after the addition of Ammonia Water, *P G*, when slightly acidulated with Hydrochloric Acid should not respond to the time-limit test for heavy metals, *U S P*

Ammonium Oxalate—An aqueous solution (1-20) after the addition of Ammonia T S should not be affected by T S of Ammonium Oxalate, *P G*

Sodium Phosphate—An aqueous solution (1-20) after the addition of Ammonia T S should not be affected by T S of Sodium Phosphate, *P G*

Barium Nitrate—An aqueous solution (1-20) should not be affected by T S of Barium Nitrate, *P G*

Silver Nitrate—An aqueous solution (1-20) should not be affected by T S of Silver Nitrate, *P G*

Potassium Ferrocyanide—20 c c of an aqueous solution (1-20) should not be rendered blue by 0.5 c c of T S of Potassium Ferrocyanide, *P G*

Chlorine and Chloroform—If to 10 c c of the aqueous solution (1-20) of the salt 1 c c of Chloroform be added and Chlorine Water introduced drop by drop with agitation, the Chloroform should not acquire a violet tint, *U S P*

Sulphuric Acid—If 0.1 gramme of the salt be sprinkled upon 1 c c of Sulphuric Acid the latter should not become coloured, *P G*, no yellow colour should appear, *U S P*

Not Official

MISTURA SALINA—Potassium Nitrate, 5 grains, Spirit of Nitrous Ether, 20 minims, Burnt Sugar, 5 minims, Camphor Water, to $\frac{1}{2}$ fl oz—*Central Throat*

Potassium Nitrate, 10 grains, Solution of Ammonium Acetate, 3 fl drms, Spirit of Nitrous Ether, 30 minims, Water to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

Potassium Citrate, $\frac{1}{2}$ drms, Spirit of Nitrous Ether, 30 minims, Solution of Acetate of Ammonium, 1 fl drms, Camphor Water, to 1 fl oz—*St Mary's*

Potassium Nitrate, 10 grains, Solution of Ammonium Acetate, 3 fl drms, Spirit of Nitrous Ether, 30 minims, Water, to $\frac{1}{2}$ fl oz—*London*

MISTURA SALINA ANODYNA—Tincture of Opium, 10 minims, Saline Mixture, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

SAL PRUNELLA—Potassium Nitrate fused and moulded into balls

CHARTA NITRATA (Austr, Belg, Dan, Fr, Ger, Ital, Norw, Port, Swed and Swiss)—Soak porous paper in a saturated solution of Nitre, and dry Roll it up and burn in a candlestick Used in asthma

The paper is sometimes impregnated also with Compound Tincture of Benzoin, Spirit of Camphor, Oils of Cassia, Cinnamon and Santal, and Tincture of Sumbul

CHARTA NITRATA ET CHLORATA—Soak porous paper in a saturated solution of Potassium Nitrate and Potassium Chlorate, and dry Used in asthma.

POTASSII PERMANGANAS.

POTASSIUM PERMANGANATE

NO Syn — KALIUM HYPERMANGANICUM

 $K_2Mn_2O_8$, eq 313 74FR, PERMANGANATE DE POTASSIUM, GER, KALIUMPERMANGANAT,
ITAL, PERMANGANATO DI POTASSIO, SPAN, PERMANGANATO POTASICO

Slender, dark, purple, odourless, prismatic crystals, possessing a purplish-blue metallic lustre, and a peculiar characteristic taste, at first sweet and afterwards unpleasant and somewhat astringent

It should be kept in well-stoppered bottles and protected as far as possible from the light and from dust. When pure it is a permanent salt

It should be handled with caution as when brought into contact with easily oxidisable substances, *e.g.*, Alcohol, Gallic and Tannic Acid, Glycerin, essential Oils, etc., it readily parts with its Oxygen, the action being very violent and frequently attended by explosion. Its solutions when mixed with Hydrogen Peroxide evolve nascent Oxygen

Solubility.—1 in 18 of Water, 1 in 3 of boiling Water

Medicinal Properties.—A powerful deodorant, a weak antiseptic. Useful internally in amenorrhœa. Externally, as a wash for foul ulcers and chancres and in ozœna, as an antiseptic gargle in throat affections

In snake bites, Lauder Brunton recommends that the wound be scraped with a clean knife, and then powdered crystals of Potassium Permanganate rubbed into the wound

Weak solution (1 in 2000) injected in gonorrhœa — *B M J E* '95, i 60, *M P* '95, i 431, *B M J E* '89, ii 88

A 1 in 2000 solution as a douche in vulvo-vaginitis in children — *Pr* lxxiv 225

Case of poisoning by repeated small doses (about 2 grains). Recovery — *L* '99, ii 1468

Found always a valuable remedy in snake bites if given in time — *L* '02, ii 1711, '03, i 139, *P J* '03, i 13, *L* '05, ii 609

In certain forms of menstrual suffering. Striking and permanent results obtained by its use — *L* '02, ii 1757

In asylum dysentery, the lower bowel washed out night and morning with a weak solution (2 to 4 grains to the pint) — *L* '02, i 588

In lupus, painted with a saturated solution, or dusting with powdered Permanganate — *L* '03, i 55, *B M J* '03, ii 194

Dose.—1 to 3 grains = 0.06 to 0.2 grammes

Prescribing Notes.—*It can be made into a pill with Massa Paraffini. It is not given in solution on account of its disagreeable taste*

It is the practice to coat Permanganate pills with Sandarach varnish, but the Alcohol contained in the varnish is liable to be oxidised at the expense of the Permanganate

Incompatibles.—Animal or vegetable matters, and any reducing agent
It is also incompatible with Morphine

Official Preparation.—Liquor Potassii Permanganatis

Not Official.—Gargarisma Potassii Permanganatis, Caloni Permanganas and Sodii Permanganas

Foreign Pharmacopœias.—Official in U.S., Austr (Kalium Hyper-crystallisatum), Belg (Kalium Permanganicum),
Ned (Hypermanganas Kalicus), Dutch (Permanganas Kalii)

ganas Kalicus), Fr (Permanganate de Potassium), Ger and Jap (Kalium Permanganicum), Hung, Russ and Swiss (Kalium Hypermanganicum), Ital (Permanganato di Potassio), Mex (Permanganato de Potasio), Port (Permanganato de Potassa), Span (Permanganato Potasico), Ital has also a crude salt for disinfecting purposes

Tests—Potassium Permanganate when heated decrepitates, and when more strongly heated it decomposes, with the evolution of Oxygen, leaving a residue of Potassium Manganate and Manganese Dioxide. It dissolves in Water, forming a deep purple coloured solution which is neutral in reaction towards Litmus paper. The aqueous solution of the residue obtained on strongly heating the salt is alkaline in reaction towards red Litmus paper. When heated with a mixture of Alcohol, Water and Sulphuric Acid it evolves an odour of Acetaldehyde, the purple-coloured solution becoming colourless, and it then affords the tests distinctive of Potassium given under that heading. The addition of Oxalic Acid Solution, Ferrous Sulphate Solution or Hydrogen Dioxide Solution to an acidified aqueous solution immediately decolorises it. It is officially required to yield 97.9 p.c. of pure Potassium Permanganate, as volumetrically determined by titration with Normal Volumetric Oxalic Acid Solution as described in the small type below. The *USP* requires that it should contain not less than 99 p.c. of pure Potassium Permanganate as volumetrically determined by titration with Tenth-normal Volumetric Oxalic Acid Solution as described below. The *PG* does not include a method of determination.

The more generally occurring impurities are Aluminium, Ammonium, Calcium, Iron, Lead, Magnesium and Sodium, Carbonates, Chlorides, Nitrates and Sulphates. In testing for these impurities a solution of the salt should be treated with sufficient Alcohol to cause complete decolorisation and the liquid filtered, a portion of the filtered liquid should neither afford a turbidity nor a flocculent precipitate when boiled with Ammonia Solution, it should not evolve an odour of Ammonia when boiled with Potassium Hydroxide Solution, when acidified with diluted Hydrochloric Acid it should afford no darkening in colour on the addition of Hydrogen Sulphide Solution. When treated with Ammonium Chloride Solution and Sodium Phosphate Solution it should afford neither a distinct turbidity nor a precipitate. A solution of the salt should not possess a decidedly alkaline reaction to red Litmus paper. A portion of the solution decolorised by Alcohol should afford no reaction when examined by the Silver Nitrate, Barium Chloride or Diphenylamine and Sulphuric Acid tests described below. In testing for Nitrates the *PG* decolorises the Potassium Permanganate with Oxalic Acid and examines the filtrate with the Sulphuric Acid and Ferrous Sulphate test described in the small type paragraph.

Litmus—An aqueous solution is neutral to Litmus, *PG* and *USP*.

For applying tests of purity the *PG* and *USP* direct the preparation of a solution as follows—0.5 gramme of the salt is boiled with 2 c.c. of Alcohol and 25 c.c. of Water and the liquid filtered. The *USP* directs 4 c.c. of Alcohol, 20 c.c. of Water, and boiling until the salt is completely decomposed. The filtrate is clear and colourless.

Barium Chloride or Nitrate—5 c c of the above filtrate acidulated with Nitric Acid should not be rendered more than very slightly turbid by T S of Barium Chloride, *U S P*, not more than opalescent by T S of Barium Nitrate, *P G*

Silver Nitrate—Another portion of the filtrate acidulated with Nitric Acid should give with Silver Nitrate T S no precipitate or cloudiness, *U S P*, should not be rendered more than opalescent, *P G*

Diphenylamine and Sulphuric Acid—If to another portion of 5 c c of the filtrate 1 drop of Diphenylamine T S be added, and then 1 c c of Sulphuric Acid be introduced so as to form a layer beneath, no blue colour should appear at the line of contact, *U S P*

Oxalic Acid, Sulphuric Acid and Ferrous Sulphate—If Oxalic Acid be gradually added to a solution of 0.5 gramme of the salt in 5 c c of hot Water until decolorisation occurs and then filtered, 2 c c of the filtrate mixed with 2 c c of Sulphuric Acid and 1 c c of Ferrous Sulphate T S poured on as a layer should show a very coloured zone, *P G*

Volumetric Determination—1 gramme of the salt dissolved in Water, and mixed with 5 c c of diluted Sulphuric Acid should require 31.2 c c of Normal Oxalic Acid Volumetric Solution for complete decolorisation, *B P*. The *U S P* directs 0.1 gramme of the salt to be dissolved in 100 c c of Water to which 1 c c of Sulphuric Acid and 35 c c of Tenth normal Volumetric Solution of Oxalic Acid have been previously added, when not more than 3.5 c c of Tenth normal Volumetric Potassium Permanganate Solution should be required to impart a permanent pink tint

Preparation

LIQUOR POTASSII PERMANGANATIS. SOLUTION OF POTASSIUM PERMANGANATE

Dissolve 87½ grains of Potassium Permanganate in Distilled Water, *q s* to yield 20 fl oz (1 in 100)

Dose.—2 to 4 fl drm = 7.1 to 14.2 c c

110 minims contain 1 grain

It needs filtration, glass-wool is best for the purpose

Diluted with 40 to 80 parts of Water, is useful as a gargle, or as a cleansing wash for foul ulcers, etc

Foreign Pharmacopœias—Official in Mex, 1 in 500 Not in the others

Not Official

GARGARISMA POTASSII PERMANGANATIS—Solution of Potassium Permanganate, 12 minims, Water, to 1 fl oz—*St Mary's, University and Westminster*

Solution of Potassium Permanganate, ½ fl oz, Water, *q s* to make 20 fl oz—*St Thomas's*

The deodorant and disinfectant properties of this gargle may be increased by the addition of 2 minims of diluted Sulphuric Acid per fl oz—*St Thomas's*

This has been incorporated in the *B P C*

Solution of Potassium Permanganate, 10 minims, Water, to 1 fl oz—*London*

Potassium Permanganate, ½ grain, Water, to 1 fl oz—*Throat*

CALCII PERMANGANAS—Calcium Permanganate is very soluble in Water. It is stated to be less nauseous than Potassium Permanganate, but there is not much difference between them in this respect

Strichnine Sulphate and Hydrochloride, Morphine, and Aconitine have been shown to yield an innocuous product when treated with a 5 p c solution of this salt—*J C S 1905, Abs 1107*

Sodii Permanganas in solution is used as a disinfectant. It is so soluble in Water that it is difficult to crystallise

POTASSII SULPHAS.

POTASSIUM SULPHATE

 K_2SO_4 , eq 173.00FR, SULFATE NEUTRE DE POTASSIUM, GFR, KALIUMSULFAT,
ITAL, SOLFATO DI POTASSIO, SPAN, SULFATO POTASICO

Hard, transparent, colourless, six-sided, rhombic prisms, terminated by pyramids, or a white, odourless powder, having a somewhat bitter saline taste Permanent in the air

Potassium Sulphate was long known as **Sal Polychrestum**, and the Bisulphate (the residue from making Nitric Acid) as **Sal Enixum**

In Scotland, Sal Polychrestum means Sulphas Potassæ c Sulphure — *Ph Edinburgh*

Solubility — 1 in 10 of cold Water, 1 in 4 of boiling Water Insoluble in Alcohol (90 p c)

Medicinal Properties — Mild, saline cathartic, usually operating without irritation if given well diluted Generally given in combination with Rhubarb A useful purgative in hepatic sluggishness

Dose — 10 to 40 grains = 0.65 to 2.6 grammes

Official Preparations — Used in the preparation of *Pilula Colocynthis Composita* and *Pulvis Ipecacuanhæ Compositus* Contained in *Pilula Colocynthis et Hyoscyami*, and *Pilula Ipecacuanhæ cum Scilla*

Foreign Pharmacopœias — Official in U.S., Dan., Dutch, Norw. and Swed. (*Sulphas Kalicus*), Fr. (*Sulfate Neutre de Potassium*), Belg., Ger., Hung., Jap., Russ. and Swiss (*Kalium Sulfuricum*), Mex. (*Sulfato de Potasio*), Port. (*Sulfato de Potassa*), Span. (*Sulfato Potasico*) Not in Austr.

Tests — Potassium Sulphate when heated decrepitates and fuses at a bright red heat It dissolves in Water, forming a clear solution which should be neutral in reaction towards Litmus paper and which should yield the tests distinctive of Potassium given under that heading, and which should yield with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid It is officially required to yield 99.9 p c of pure Potassium Sulphate as gravimetrically determined by precipitation as Barium Sulphate, the washed and dried precipitate from a solution of 1 gramme of the salt in Water, acidulated with Hydrochloric Acid, is required to weigh 1.339 grammes The *USP* states that it should contain not less than 99 p c of pure Potassium Sulphate, but gives no method of determination The *PG* gives neither a requisite percentage of Sulphate nor a method of determination

The more generally occurring impurities are Arsenic, Calcium, Copper, Iron, Lead, Magnesium, Sodium and Zinc The *BP* includes also Aluminium and Ammonium Chlorides and Nitrates are also likely impurities, as also Acid Potassium Sulphate Arsenic, if present, may be detected by the modified Gutzert's test Iron, Copper, Lead and Zinc, if present, are indicated by the Hydrogen Sulphide test described below, the *PG* includes a separate test for Iron with Potassium Ferrocyanide Solution, Calcium and Magnesium by the Ammonium Oxalate

and Sodium Phosphate tests described in small type Aluminium may be detected by the turbidity or flocculent precipitate produced on boiling the solution with Ammonia Solution Ammonium salts by the evolution of Ammonia gas when a solution of the sample is boiled with Liquor Potassæ Chlorides, if present, may be detected by the Silver Nitrate test described in small type Acid Potassium Sulphate, if present, is indicated by the behaviour of the aqueous solution of the specimen towards Litmus paper, solutions of the Acid Sulphate have a decidedly acid reaction towards blue Litmus paper A solution of the salt, when carefully mixed with Sulphuric Acid, the mixture being kept cool, should afford no brown ring at the junction of the two liquids when a Solution of Ferrous Sulphate is carefully poured upon its surface

Hydrogen Sulphide—The aqueous solution (1-20) should not be affected by T S of Hydrogen slightly acidulated with Hydrochloric Acid should not respond test for heavy metals—*U S P*

Gutzeit's Test—5 c c of an aqueous solution of the salt (1-10) should not respond to the modified Gutzeit's test for Arsenic, *U S P*

Ammonium Oxalate—The aqueous solution (1-20) of the salt should not be affected by T S of Ammonium Oxalate, *P G*

Silver Nitrate—The aqueous solution (1-20) of the salt should not be affected by T S of Silver Nitrate, *P G*

Sodium Phosphate—The aqueous solution (1-20) of the salt should not be affected by T S of Sodium Phosphate, *P G*

Potassium Ferrocyanide—20 c c of an aqueous (1-20) solution of the salt should not be rendered blue by 0.5 c c of T S of Potassium Ferrocyanide, *P G*

POTASSII TARTRAS.

POTASSIUM TARTRATE

$K_2C_4H_4O_6$, H_2O , eq 242.46

FR, TARTRAT DE POTASSE NEUTRE, GER, KALIUMTARTRAT, ITAL, TARTRATO NEUTRO DI POTASSIO, SPAN, TARTRATO POTASICO NEUTRO

Colourless, translucent, prismatic crystals, or as a white, crystalline, slightly deliquescent powder, having a saline and bitter taste. It may be obtained by neutralising the Acid Tartrate with Potassium Carbonate

Solubility.—10 in 6 of Water Insoluble in Alcohol (90 p c)

Medicinal Properties.—A mild laxative, acting without much pain, and producing doses, antacid, diuretic and alterative

Dose.—30 to 240 grains = 2 to 16 grammes

Foreign Pharmacopœias—Official in Dan and Norw (Tartras Kalicus), Ger, Hung, Jap and Russ (Kalium Tartaricum), Ital (Tartrato Neutro di Potassio), Mex (Tartrato de Potasio Neutro), Port (Tartarato de Potassa), Span (Tartrato Potasico Neutro). Not in Austr, Belg, Dutch, Fr, Swiss or U S

Tests—Potassium Tartrate when heated decomposes, evolving an odour of burnt sugar and leaving behind a black residue, the aqueous solution of which possesses an alkaline reaction towards red Litmus paper. It dissolves readily in Water, forming a solution which has a slightly alkaline reaction towards red Litmus paper. When acidified with Hydrochloric Acid it should yield the tests distinctive of Potassium given under that heading. The aqueous solution yields on the addition of Calcium Chloride Solution in excess a white granular precipitate soluble when freshly precipitated in cold, moderately concentrated Potassium Hydroxide Solution, reprecipitated on boiling, it is soluble also in diluted acids. Silver Nitrate Solution affords a white precipitate soluble in diluted Nitric Acid and in Ammonia Solution, the solution obtained by the use of the latter reagent, when boiled in a perfectly clean test-tube, deposits a mirror of metallic Silver on the sides of the tube. When mixed with Ferrous Sulphate Solution, a few drops of Hydrogen Peroxide Solution and an excess of Potassium Hydroxide Solution, a purple or violet coloration is produced. It is officially required to contain 101.8 p.c. of pure Potassium Tartrate as volumetrically determined by the direct titration of the solution of the residue left on ignition of the dried salt, with Volumetric Sulphuric Acid Solution, the inaccuracy being due to the incorrectness of the official formula. The salt is official in the *P.G.*, but no requisite percentage of pure Tartrate is indicated, nor is a method of determination given.

The more generally occurring impurities are Calcium, Copper, Iron and Lead, Magnesium, Sodium, Chlorides, Sulphates and Acid Potassium Tartrate. Calcium, if present, may be detected by Ammonium Oxalate test after separation of the greater portion of the Potassium Tartrate as insoluble Acid Tartrate of Potash by precipitation with diluted Acetic Acid. Copper, Lead, and Iron may be detected by the Hydrogen Sulphide test described below, the *P.G.* includes a separate test for Iron with Potassium Ferrocyanide Solution, Magnesium by Sodium Phosphate Solution, Chlorides and Sulphates may be detected by the Barium Nitrate and Silver Nitrate tests described in the small type paragraphs. Potassium Acid Tartrate, if present, may be detected by the behaviour of the aqueous solution towards blue Litmus paper, Acid Potassium Tartrate having a decidedly acid reaction. The salt should not evolve an odour of Ammonia when warmed with Sodium Hydroxide Solution, when acidified with Hydrochloric Acid and inserted on a loop of platinum wire into a non-luminous flame it should not colour the flame a distinct or permanent yellow colour.

Acetic Acid and Ammonium Oxalate—If 1 gramme of Potassium Tartrate be dissolved in 10 c.c. of Water and the solution agitated with 5 c.c. of Diluted Acetic Acid a crystalline separation occurs. The liquid poured off and diluted with an equal part of Water should not be affected by 8 drops of Ammonium Oxalate T.S. within 1 minute, *P.G.*

Hydrogen Sulphide—The aqueous solution (1-20) should not be affected by T.S. of Hydrogen Sulphide, *P.G.*

Barium Nitrate—The aqueous solution (1-20), after the addition of Nitric

Acid and the removal of the crystalline precipitate formed, should not be affected by T S of Barium Nitrate, *P G*

Silver Nitrate—Another portion of an aqueous solution treated as above should not be rendered more than opalescent by T S of Silver Nitrate, *P G*

Potassium Ferrocyanide—20 c c of an aqueous solution (1-20) should not be rendered blue by 0.5 c c of T S of Potassium Ferrocyanide, *P G*

Volumetric Determination—The filtered aqueous solution obtained by dissolving in Water, the alkaline residue left on exposing 1 gramme of the salt to a red heat should neutralise 8.4 c c of Volumetric Sulphuric Acid Solution, *B P*

POTASSII TARTRAS ACIDUS.

ACID POTASSIUM TARTRATE

B P Syn—BITARTRATE OF POTASSIUM, PURIFIED CREAM OF TARTAR

N O Syn—TARTARUS DEPURATUS KALIUM HYDROTARTARICUM

$\text{KC}_4\text{H}_5\text{O}_6$, eq 186.75

Fr, TARTRATE ACIDE DE POTASSIUM, Ger, WEINSTEIN, Ital, TARTRATO ACIDO DI POTASSIO, Span, BITARTRATO POTASICO

Colourless or slightly opaque rhombic crystals, or masses of crystals, or as a white, crystalline, gritty powder, permanent in the air. Its chief source is the crude Cream of Tartar or Argol, deposited during vinous fermentation.

It should be kept in well-stoppered glass bottles.

Solubility—1 in 200 of cold Water, 1 in 16 of boiling Water. Insoluble in Alcohol (90 p c)

Medicinal Properties.—Cathartic, diuretic and refrigerant. Much used in febrile and diopsical affections, in chronic cardiac and hepatic diseases, combined with Sulphur it is a useful laxative in hæmorrhoids.

‘Imperial Drink’ for patients suffering from the thirst accompanying chronic Bright’s disease, Acid Tartrate, 1 drm, Saccharin, 1 grain, Lemon Oil, 8 minims, boiling Water, to 1 pint.—*P*, lxvii 656

Dose.—20 to 60 grains = 1.3 to 4 grammes

Official Preparations—Contained in Confectio Sulphuris, Trochiscus Sulphuris, and Pulvis Jalapæ — 1 f Acidum Tartaricum, Antimonium , , Tartarus, and Soda Tartarata

Not Official—Soluble Cream of Tartar

Foreign Pharmacopœias—Official in all, Austr and Hung (Kalium hydrotartaricum), Belg (Bitartras Potassæ depuratus), Dan, Norw and Swed (Bitartras Kalicus), Dutch (Tartras Kalicus Acidus), Fr (Tartrate Acide de Potassium), Ger and Swiss (Tartarus depuratus) Ital (Tartrato Acido di Potassio), Jap (Kalium Bitartaricum), Mex (Tartrato de Potasio Acido), Port (Bitartrato de Potassa), Russ (Kalium Bitartaricum depuratum and Purum, Span (Bitartrato Potasico), U S (Potassii Bitartras)

Tests—Potassium Acid Tartrate when strongly heated chars and emits inflammable vapours. When still more strongly heated it leaves

a blackish residue, which, when dissolved in Water, filtered and neutralised with Hydrochloric Acid, affords the tests distinctive of Potassium given under that heading. When neutralised with Potassium or Sodium Hydroxide Solution it yields, on the addition of Calcium Chloride Solution in excess, a white granular precipitate soluble when freshly precipitated in cold, moderately concentrated Potassium Hydroxide Solution, reprecipitated on boiling. In a similar solution Silver Nitrate Solution yields a white precipitate soluble in Nitric Acid and in Ammonia Solution, its solution in the latter reagent, when boiled in a perfectly clean test-tube, affords a mirror of metallic Silver on the sides of the tube. It is slightly soluble in Water, the solution being acid in reaction towards blue Litmus paper. It is officially required to contain 97.1 p.c. of pure Potassium Hydrogen Tartrate as volumetrically determined by direct titration of the salt with Volumetric Sodium Hydroxide Solution as indicated below. The direct titration of a sample should be supplemented by a determination of the alkalinity of the soluble ash. A sample containing a judicious proportion of Potassium Acid Sulphate might pass the direct titration test, but would show a reduced alkalinity of the ash. In a pure sample the amount of Volumetric Sodium Hydroxide Solution required by direct titration should be equal to the amount of Volumetric Acid Solution required to neutralise the soluble ash, working on the same weight of substance in each case. The *U.S.P.* requires the salt to contain not less than 99 p.c. of pure Potassium Bitartrate as volumetrically determined by titrating with Semi-normal Volumetric Sulphuric Acid Solution, the solution obtained by extracting the residue left after ignition with Water, using Methyl Orange Solution as an indicator.

The more generally occurring impurities are Copper and Iron, with the tests for which it is officially required to show no characteristic reaction, Calcium, Magnesium, Sodium, Chlorides or Sulphates, with the tests for which it is officially required to show only the slightest reaction. The term 'slightest reaction' in this instance appears to be defined by the total limit of 2½ p.c. on the dried salt, the official requirement being that this amount of impurity should not be exceeded. Lead, Copper and Iron, if present, are shown by the Hydrogen Sulphide test described below, Calcium by the Ammonium Oxalate test, Magnesium by the addition of Sodium Phosphate Solution to the filtrate from the Ammonium Oxalate. A standard has been suggested (*C.D.* '08, 1796) of 5 parts per million for Lead and 2 parts per million for Arsenic Chlorides and Sulphates, if present, are indicated by the Silver Nitrate and Barium Chloride tests given below. In addition to the above-named impurities, Starch, Kaolin, Calcium Phosphate and other insoluble matter, Ammonium salts, Aluminium and Phosphates may be present, and the *U.S.P.* includes tests for these impurities. The undermentioned test with Ammonia Water serves as a test for Starch, Kaolin, Calcium Phosphate and other insoluble matter. The presence of Ammonium salts is indicated by the behaviour of the solution when examined by the tests with Potassium or Sodium

Test Solution, Aluminium by the Potassium Hydroxide test ; and Phosphate by the Ammonium Molybdate test given below. A method for the exhaustive examination of Cream of Tartar is given (*Analyst*, '96, 174)

Ammonia—A solution of 0.5 gramme of the salt in 3 c.c. of Ammonia T.S. should leave no insoluble residue, *U S P*

Barium Nitrate—5 grammes of the salt agitated with 100 c.c. of Water and filtered, should give a filtrate which, after the addition of Nitric Acid, should not be affected by T.S. of Barium Nitrate, *P G*

Silver Nitrate—A filtrate obtained as above should not be rendered more than faintly opalescent with T.S. of Silver Nitrate after the addition of Nitric Acid, *P G*

Hydrogen Sulphide—The solution of 1 gramme of the salt in Ammonia T.S. should not be affected by T.S. of Hydrogen Sulphide, *P G*. The *U S P* requires that the filtrate of the salt, slightly acidulated with Hydrochloric Acid, should give a time-limit test for heavy metals.

Potassium or Sodium Hydroxide—If the salt be warmed with Sodium Hydroxide T.S. (15 p.c.) it should not evolve Ammonia, *P G*. The *U S P* uses Potassium Hydroxide T.S.

Ammonium Oxalate—If a mixture of 1 gramme of the salt with 5 c.c. of diluted Acetic Acid be allowed to stand for half an hour with frequent agitation and then mixed with 25 c.c. of Water, the clear liquid poured off from the deposit should show no alteration within 1 minute on the addition of 8 drops of T.S. of Ammonium Oxalate, *P G*

Potassium Hydroxide Solution after Ignition—If 1 gramme of Potassium Bitartrate be well triturated with about 1 gramme of Potassium Carbonate and 0.5 gramme of Potassium Nitrate, and the mixture heated gradually to dull redness in a porcelain crucible, and if, upon being treated with a slight excess of diluted Hydrochloric Acid, the filtrate, upon being made slightly alkaline with Potassium Hydroxide T.S., should not yield a gelatinous precipitate soluble in excess of the reagent (absence of Alum), *U S P*

Ammonium Molybdate Solution—If a precipitate be obtained which is insoluble it should be collected and thoroughly washed with distilled Water, and dissolved in hot diluted Nitric Acid, the addition of an excess of Ammonium Molybdate T.S. to this solution should not produce a yellow precipitate (absence of Phosphates), *U S P*

Volumetric Determination—Not less than 5.2 c.c. of Sodium Hydroxide Volumetric Solution should be necessary to neutralise 1 gramme of the salt, *B P*. The *U S P* directs that 1 gramme of the salt be thoroughly carbonised at a temperature not exceeding red heat, and the residue extracted with boiling Distilled Water until the washings cease to react with Methyl Orange T.S. The mixed filtrate or washings should require for complete neutralisation not less than 10.6 c.c. of Semi-normal Volumetric Sulphuric Acid Solution, Methyl Orange T.S. being used as indicator.

Not Official

TARTARUS BORAXATUS TARTRATE BORICO-POTASSIQUE. SOLUBLE CREAM OF TARTAR—Soluble Cream of Tartar is a white, amorphous powder soluble in its own weight of Water. The proportions are—Dan., Norw. and Swed., Potassium Tartrate 2, Borax 1, Dutch and Ger., Potassium Acid Tartrate 5, Borax 2, dissolve the Borax and the Acid Tartrate in Water by the aid of heat, and evaporate to dryness, Ital. (Tartrato Borico-Potassico), proportions not given, Span., Potassium Acid Tartrate 4, Boric Acid 1, Fr. (Borotartrate de Potassium), and Mex. (Tartrato borico-potassico), Potassium Bicarbonate 10, Tartaric Acid 10, Boric Acid 5, Port., with Boric Acid and Potassium Acid Tartrate, but no quantities given.

PRUNI VIRGINIANÆ CORTEX.

VIRGINIAN PRUNE BARK

WILD CHERRY—U S P

The Bark of *Prunus serotina*, Ehrh, it is believed to be stronger in the autumn than in the spring, and is required to be collected in the autumn both by *B P* and *U S P*

In addition to astringent Tannins, this bark contains **Amygdalin** and **Emulsin**, which on treatment with Water develop Hydrocyanic Acid (in a manner similar to the Cherry Laurel), to which the sedative effects of its preparations are probably due

Medicinal Properties—Sedative Highly useful in full doses for resultless hacking cough in phthisis and chronic bronchitis The Syrup is also useful as a flavouring vehicle for nauseous medicines

Official Preparations—Syrupus Pruni Virginianæ, and Tinctura Pruni Virginianæ

Not Official—Fluidextractum Pruni Virginianæ, Infusum Pruni Virginianæ

Foreign Pharmacopœias—Official in U S Not in the others U S has also an Infusum and Fluid Extract

Descriptive Notes—This bark was formerly supposed to be derived from *Prunus Virginiana*, Mill, whence the official name, it is now officially referred to *Prunus serotina*, Ehrh. The bark varies much in appearance, according to the age of the tree, the young bark being thin, with a reddish external thin papery layer $\frac{1}{10}$ to $\frac{1}{15}$ in (1.5 to 2.5 mm) in thickness, with an underlying greenish layer marked with transverse lenticular growths, but in older bark the surface is rough and darker brown and thicker, in both the fracture is short and granular, and the under surface exhibits a minute porous network with small fissures. It has an astringent, bitter taste, and a slight almond flavour when chewed. The young bark is richer in Hydrocyanic Acid, but the *B P* does not exclude old bark. Both the *B P* and *U S P* state that the bark should be collected in autumn, but the *U S P* describes only the young bark, in curved pieces 3 to 7 cm long and 0.5 to 4.0 mm in thickness. Young bark about $\frac{1}{12}$ in (2 mm) is the best, when $\frac{1}{2}$ to $\frac{1}{2}$ in (3 to 4 mm) in thickness it is likely to be less active. The bark is apt to deteriorate if kept long. The bark of other species of *Prunus* is sometimes substituted for that of *Prunus serotina*. One of these, described by Finckh, has a fibrous fracture and gives a darker coloured syrup and is devoid of the characteristic odour and taste. Under the microscope the leading features of the powder are the irregularly shaped stone cells, the brown medullary rays 4 cells broad, large rhomboidal raphides near the stone cells, and smaller ones in the pith, sphaeraphides occur in rows in the soft bast and are coloured brown by solution of Caustic Potash. The tissue also contains an abundance of small Starch grains and brown particles of a kind of Tannin which turn black with Ferric salts. In the false bark long bast fibres are present, but no stone cells of irregular shape.

Tests.—Virginian Prune Bark contains from 3 to 4 p c of ash, and should not exceed 6 p c

Preparations

SYRUPUS PRUNI VIRGINIANÆ. SYRUP OF VIRGINIAN PRUNE

Percolate 3 of Virginian Prune Bark, in No 20 powder, with Distilled Water to yield 9, Refined Sugar, in coarse powder, 15, Glycerin, 1½. Dissolve 15 of Refined Sugar in the liquid, by agitation, without heat, add 1½ of Glycerin, strain, pour Distilled Water over the strainer to make up to 20

Dose.—½ to 1 fl dm = 1 8 to 3 6 c c

Foreign Pharmacopœias—Official in U S, Wild Cherry 15, Sugar 70, Glycerin 15, Water to make 100

Tests.—Syrup of Virginian Prune has a sp gr of 1 300 to 1 310

TINCTURA PRUNI VIRGINIANÆ. TINCTURE OF VIRGINIAN PRUNE

Macerate 4 of Virginian Prune Bark, in No 20 powder, with 7½ of Distilled Water for 24 hours, add 12½ of Alcohol (90 p c) and continue the maceration for 7 days

Dose.—½ to 1 fl drm = 1 8 to 3 6 c c

Tests.—Tincture of Virginian Prune has a sp gr of 0·935 to 0 940, contains about 3 p c w/v of total solids and about 54 p c w/v of Absolute Alcohol

Not Official

FLUIDEXTRACTUM PRUNI VIRGINIANÆ—Wild Cherry, in No 30 powder, 100, is percolated with a mixture of Glycerin 20, Alcohol (95 p c) 20, and Water 60, and then percolation is continued with a mixture of 20 of Alcohol (95 p c) and 80 of Water, until the product measures 100 **Dose**—½ to 1 fl dm (1 8 to 3 75 c c)—*U S P*

This has been incorporated in the *B P C*, but whereas the *U S P* continue the percolation to produce 100, the *B P C* percolate to exhaustion, reserving the first 90 and concentrate the subsequent weaker percolate to a soft extract and add to the reserved portion

INFUSUM PRUNI VIRGINIANÆ—Wild Cherry, in No 20 powder, 4, Glycerin 5, Water *qs* to make 100. Moisten the powder with 6 of Water, allow it to macerate for 1 hour, pack into a glass percolator, and having put the Glycerin into the bottle gradually pour Water upon the powder, and continue percolatio product measures 100, and mix well—*U S P*

This has been incorporated in the *B P C*

PRUNUM.

PRUNES

The dried ripe Fruits of *Prunus domestica* var. *Juhana*

Imported from the South of France

Medicinal Properties—Mild laxative, nutritious and demulcent. Often used in dietetic treatment of constipation as a laxative

Official Preparation—Contained in *Confectio Sennæ*

Foreign Pharmacopœias—Official in Dutch, Pruna, Mex (Ciruelo de Espana), Port (Ameixas Passadas), US Not in the others

Descriptive Notes—The dried fruits of *Prunus domestica* var *Juhana* DC are official in the *BP*, but no special variety of the species is specified in the *USP*. The size given in the *BP* is $1\frac{1}{4}$ in (3 cm) long, and in the *USP* 3 to 4 cm. Plums differ in size and shape and in the character of the stone. The stone of the var *Juhana* is described by Hanbury as $\frac{7}{16}$ to $\frac{9}{16}$ in long and $\frac{5}{16}$ to $\frac{6}{16}$ in broad, broadly rounded at the upper end and slightly mucronulate, narrowed and somewhat stalk-like at the lower end and truncate. The ventral suture is broader and thicker than the dorsal. German prunes (Zwetschen or Quetschen) derived from the var *Prunearuhana*, DC, are sometimes imported. These are rather larger and more elongated and have a thicker skin and a flatter narrower stone, pointed at either end, with the ventral suture more strongly curved than the dorsal, and the fruits are more prone to become covered with a saccharine efflorescence.

Not Official

PSYLLII SEMEN

The seeds obtained from *Plantago Psyllium*, L. They are small, about $\frac{1}{8}$ in long and about half as broad, they are boat shaped, convex on one side and concave on the other, they are a bright brown in colour, without odour, but have a disagreeable, somewhat acrid taste when fresh, which is removed by Alcohol with which some commercial specimens appear to have been treated. They are closely allied to the Ispaghul Seeds so well known in India, and, like them, are very mucilaginous. They have recently come into more general use in this country in some forms of habitual constipation.

Dose—A teaspoonful to a tablespoonful of the seeds in half a tumbler of Water at bedtime, or before the principal meal.

PTEROCARPI LIGNUM.

RED SANDERS-WOOD

BP Syn—RED SANDAL-WOOD

(SANTALUM RUBRUM, *USP*)

The Heart-wood of *Pterocarpus santalinus*, Linn f

From Madras and Ceylon. Used solely as a colouring agent.

Official Preparation—Used in the preparation of Tinctura Lavandulae Composita

Foreign Pharmacopœias—Official in Austr, Jap and Swed (Lignum Santali Rubrum), Dutch (Lignum Santalinum), Port (Sândalo Rubro), Span (Sândalo Rojo), US (Santalum Rubium). Not in Fr, Ger, Hung, Ital, Mex, Norw, Russ or Swiss.

Descriptive Notes—Red Sandal-Wood being much valued for turning purposes, the logs imported for medicinal use are chiefly derived from the tree stumps or roots. The wood is blackish-brown externally, but of a deep red colour in transverse section, with lighter zones. It is usually met with in retail commerce in the form of splintery raspings of a purplish-red colour. The colouring matter is

soluble in 90 p c spirit and in caustic alkalis, but is precipitated by acids. The wood has only a faintly astringent taste, and a slight aromatic odour when warmed.

Tests.—It contains about 1 p c of ash. It imparts a red colour to Alcohol, but not to Water.

Not Official

PULSATILLA.

The Herb of *Anemone Pulsatilla*, L., and of *Anemone pratensis*, L., collected soon after flowering.

It should be carefully preserved, and not kept longer than one year.

It contains an unstable body, *Anemone-camphor*, which occurs in tiny metallic prisms. It splits up into Anemonin and Anemonic Acid.

Medicinal Properties—Has been used in dysmenorrhœa with various results.

Has been recommended in orchitis and epididymitis, but in experiments at the Lock Hospital it was found to be valueless.—*L* '89, ii 216.

Foreign Pharmacopœias—Official in Fr. Not in the others.

Not Official

ALCOOLATURE D'ANEMONE PULSATILLE—Bruised fresh flowers and leaves of Pulsatilla, 1, Alcohol (95 p c), by weight, 1. Macerate for 8 days, press and filter.—*Fr*.

TINCTURA PULSATILLÆ—Carefully dried Herb, 1, Alcohol (60 p c), to percolate 10.

Dose.—5 to 30 minims = 0.8 to 1.8 c c.

A tincture of this strength was included in *B P C Formulary* 1901, and is now incorporated in the *B P C*.

Unless the Herb is very finely powdered, it answers better to soak Water for a day, and then add Alcohol to bring the mixture to the Alcohol (60 p c).

It is best prepared from the fresh herb, as in *Fr Codex*, see above.

LIQUOR CAULOPHYLLIN ET PULSATILLÆ, see p 349.

PYRETHRI RADIX.

PYRETHRUM ROOT

NO Syn—PELLITORY ROOT

Fr, PYRÈTHRE D'AFRIQUE, Ger, ROMISCHE BERTRAMWURZEL, Ital, PIRETRO, Span, RAIZ DE PELITRE.

The dried Root of *Inula racemosa* *Pyrethrum*, DC.

Collected chiefly in Algeria.

Medicinal Properties—It is powerfully stimulant to the salivary glands, causing a copious flow of saliva, and, on that account, is used as a masticatory in dryness of the mouth and throat. The Tincture is used on Cotton-Wool for relieving toothache, or when diluted, as a mouth-wash.

Official Preparation—Tinctura Pyrethri.

Not Official—Trochisci Pyrethri.

Foreign Pharmacopœias—Official in Austr, Fr (Pyrethre d'Afrique), Mex (Peritre de Africa), Port (Pyrethro), Span (Pelitre), and U S Not in the others

Descriptive Notes—Pyrethrum root, of Pellitory of Spain, occurs in cylindrical pieces tapering slightly towards either end, unbranched, and having sometimes at the apex a short tuft of soft woolly hairs. It is longitudinally furrowed, is of a dark brown colour and has a short fracture, exhibiting in transverse section a radiate structure, with dark resin cells in the cortex, and medullary rays, the wood is porous and yellow, and the bark is dark brown. It has a characteristic odour, and a slowly pungent and acrid taste, causing a flow of saliva. It has been adulterated with the root of *Corrigiola telephnifolia*, Poir., which, like Pyrethrum, is a product of Morocco. The root is similar in size, but paler, and exhibits in transverse section a series of horny concentric zones, but no resin ducts.

Tests—It contains from 4 to 5 p c of ash, and the latter figure should not be exceeded.

Preparation

TINCTURA PYRETHRI TINCTURE OF PYRETHRUM

Pyrethrum Root, 4, Alcohol (70 p c), *q s* to yield 20 (1 in 5)

The Tincture in *BP* '85 was made with Rectified Spirit or Alcohol (84 p c), *USP* employ Alcohol (95 p c), and *BPC* Alcohol (90 p c), subsequently altered to Alcohol (70 p c), all are 1 in 5.

Foreign Pharmacopœias—Official in Fr, Mex and U S, 1 in 5. Not in the others.

Tests—Tincture of Pyrethrum has a sp gr of 0.900 to 0.904, it contains from 1.5 to 2 p c w/v of total solids and about 68 p c w/v of Absolute Alcohol.

Not Official

TROCHISCI PYRETHRI—Contain 1 grain in each—*Throat*

Not Official

PYRETHRI FLORES

Syn—INSECT POWDER

The powder of the Flower heads, obtained in the Caucasus, from *Pyrethrum roseum*, Bieb., and *P. carneum*, Bieb., and in Dalmatia from *Pyrethrum cnerarice-folium*, Trev.

The active principle is an Ether soluble Resin, not a volatile Oil.

Foreign Pharmacopœias—Official in Mex (Peritre del Caucasos). Not in the others.

Keeps away troublesome insects.

Descriptive Notes—The Persian Insect powder of commerce is derived from the flowers of *Pyrethrum cneraricefolium*, of which three qualities are sold, named respectively 'closed,' 'half closed,' and 'open,' the closed being the most effective, if gathered when full grown. In the closed flowers the yellowish-white, lanceolate, acute, hairy phyllaries or outer bracts are incurved, in the half closed the Flower-heads are frequently more or less deprived of the white ligulate florets, and in the open flowers even the tubular florets may be partially fallen away. The ovary has no pappus, but the calyx forms a short membranous ring.

with 5 minute teeth, and the fruit has 5 slender rib-like wings. The powdered flowers are characterised by the epidermal papillæ of the ligulate florets being more conical and narrower, and having a thinner apex than those of *Pyrethrum roseum*, the Flower-heads of which are now seldom used, they are distinguished by the red ligulate florets, and the phyllaries with brownish-black margins. As a rule the powder is more active the more abundant the pollen and the smaller the quantity of stem tissue. See also *P J* (4) iv pp 505-507

This powder has been adulterated with the flowers of other species of *Chrysanthemum*, with White Hellebore, and with chrome yellow and with powdered Surinach leaves. See *Vogl, Pharmacognosie*, pp 116, 117

TINCTURA PYRETHRI FLORUM—The Flower-heads, in powder, 1, Alcohol (60 p c) to percolate 4

Dilute 1 to 10 of Water forms a lotion to keep away insects

This has been incorporated in *B P C*

Not Official

PYRIDIN.

C_5H_5N , eq 78.49

A colourless, volatile liquid, with a powerful and a peculiar odour. Its aqueous solution has a strong alkaline reaction to Litmus. It yields a crystalline deliquescent salt with Hydrochloric Acid.

It is a base obtained from the products of the destructive distillation of bones.

Commercially it always contains Picoline. In its cruder forms it is employed in Germany for 'denaturing' Alcohol, corresponding to 'Methylating' in this country.

Solubility—It is miscible with Water, Alcohol (90 p c), Ether, and the fixed Oils.

Medicinal Properties—Useful in the treatment of asthma, 4 or 5 grammes (62 to 77 grains) are allowed to evaporate from a flat dish in a small room, the patient being exposed to its vapour for about half an hour 3 times a day. It is most beneficial in cardiac dyspnoea, emphysema, and angina pectoris.

If the vapour be inhaled in quantity, it produces headache.

Like Nicotine, it is a good insecticide.

Official in Fr, Mex and Spain

Tests—Pyridin has a sp gr of about 0.980. It boils at about 116° C (240.8° F). When pure it has a strong alkaline reaction towards red Litmus paper but no action on Phenolphthalein Solution. When added to Copper Sulphate Solution it gives a bluish-green precipitate, soluble in excess to a dark blue liquid similar to that produced by Ammonia. It may be titrated with Normal or Tenth normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Methyl Orange solution as an indicator of neutrality. 1 cc of Pyridin requires about 12.4 cc of Normal Hydrochloric Acid. Each cc of Normal Volumetric Hydrochloric Acid Solution represents 0.07849 gramme of absolute Pyridin. The sample should not reddens Phenolphthalein Solution, indicating the absence of Ammonia. It should have little or no action on Potassium Permanganate Solution, indicating the absence of readily oxidisable organic impurities, a 0.5 p c solution of Pyridin should give a crystalline precipitate, becoming almost semi-solid when mixed with an equal volume of saturated Picric Acid Solution. It should be completely volatilised by heat without leaving any weighable residue.

PYROXYLINUM.**PYROXYLIN**

NO Syn—**Gossipium Fulminans, Fulmi Cotton, Lana Colloidi, Colloxylinum**

Fr, Fulmi Cotton, **Ger**, Kollodiumwolle, **Ital**, Cottoni Colloidio, **Span**, Piroxilina

A white, fibrous substance, having very much the appearance of ordinary cotton, but rapidly burning away with a flash when ignited. It requires to be kept in small quantities in a cool dry place, and away from lights

Pyroxylin is Dinitrocellulose $C_6H_7(NO_2)_2O$, Gun Cotton is Trinitrocellulose $C_6H(NO_2)_3O$, and is not soluble in any mixture of Alcohol and Ether

It sometimes decomposes on keeping, with disengagement of Nitrous fumes, and becomes insoluble

The safest and best plan for its preservation is to moisten the dry material with an equal weight of Methylated Spirit, and preserve in a well stoppered jar, when required for use it is quickly and easily dried—*P J* '96, ii 110, *CD* '96, ii 207

Pyroxylin is officially prepared by nitrating Cotton, 1 of Cotton being immersed in a mixture of 5 of Sulphuric Acid and 5 of Nitric Acid, the mixture being stirred during 3 minutes and the free acid separated by washing with Water till the washings no longer have an acid reaction. The Pyroxylin is drained and dried at a low temperature

The proportions vary considerably in the different Pharmacopœias (*see below*)

Solubility—Readily soluble in a mixture of equal volumes of Ether and Alcohol (90 p c), also in Acetone

Different samples of Pyroxylin vary considerably in the extent to which they are soluble

The use of Acetone as a cheap and efficient solvent for Pyroxylin was suggested in the 16th edition of the *Companion*. A good Pyroxylin will dissolve readily to the extent of 10 p c. A formula corresponding with the *BP* Collodion is Pyroxylin 1, Acetone 48, but this produces an inconveniently thin solution, and a preferable strength is Pyroxylin 3.5, Acetone *q s* to make 100. The suggestion for its use has been followed by the *BP C* with certain additions to the formula to make it more closely resemble a well known proprietary article, it appears under the title **Collodium Acetonum** as follows—Pyroxylin 5, Oil of Cloves 2, Amyl Acetate 25, Benzol 20, Acetone *q s* to produce 100. A more descriptive title for this would have been **Collodium Acetonum Compositum**, a simple solution in Acetone having been used as Acetone Collodion

Foreign Pharmacopœias—Official in Belg (**Pyroxylinum**), Dutch, Ger and Jap, Nitric Acid 400, Sulphuric Acid 1000, Purified Cotton 55, Russ uses same formula, but Purified Cotton 50, Fr (**Fulmi cotton**), Mex (**Piroxilina**), Cotton 1, Nitre 20, Sulphuric Acid 30, Port (**Algoda Polvora**), and Span (**Piroxilina**), Cotton 1, Nitre 20, Pure Sulphuric Acid (sp gr 1.84) 80, Austr, Dan, Hung, Ital, Norw, Swed, Swiss and US, no formula given

Official Preparations—Used in the preparation of Collodium, and Collodium Vesicans. Of Collodium, Collodium Flexile

Not Official.—Collodium Stypticum, Hæmostatic Collodion, Collodium Anodynum, Celloidin, Photoxylin

with 5 minute teeth, and the fruit has 5 slender rib-like wings. The powdered flowers are characterised by the epidermal papillæ of the ligulate florets being more conical and narrower, and having a thinner apex than those of *Pyrethrum roseum*, the Flower-heads of which are now seldom used, they are distinguished by the red ligulate florets, and the phyllaries with brownish-black margins. As a rule the powder is more active the more abundant the pollen and the smaller the quantity of stem tissue. See also *P J* (4) iv pp 505-507.

This powder has been adulterated with the flowers of other species of *Chrysanthemum*, with White Hellebore, and with chrome yellow and with powdered Sumach leaves. See *Vogl, Pharmacognosie*, pp 116, 117.

TINCTURA PYRETHRI FLORUM—The Flower-heads, in powder, 1, Alcohol (60 p c), to percolate 4.

Diluted 1 to 10 of Water forms a lotion to keep away insects.

This has been incorporated in *B P C*.

Not Official

PYRIDIN

C_5H_5N , eq 78.49

A colourless, volatile liquid, with a powerful and a peculiar odour. Its aqueous solution has a strong alkaline reaction to Litmus. It yields a crystalline chloride salt with Hydrochloric Acid.

It is a base obtained from the products of the destructive distillation of bones,

Commercial Picoline. In its cruder forms it is employed in Germany for 'col', corresponding to 'Methylating' in this country.

Solubility—It is miscible with Water, Alcohol (90 p c), Ether, and the fixed Oils.

Medicinal Properties—Useful in the treatment of asthma, 4 or 5 grammes (62 to 77 grains) are allowed to evaporate from a flat dish in a small room, the patient being exposed to its vapour for about half an hour 3 times a day. Is most beneficial in cardiac dyspnoea, emphysema, and angina pectoris.

If the vapour be inhaled in quantity, it produces headache.

Like Nicotine, it is a good insecticide.

Official in Fr, Mex and Span

Tests—Pyridin has a sp gr of about 0.980. It boils at about 116° C (240.8° F). When pure it has a strong alkaline reaction towards red Litmus paper, but no action on Phenolphthalein Solution. When added to Copper Sulphate Solution it gives a bluish-green precipitate, soluble in excess to a dark blue liquid similar to that produced by Ammonia. It may be titrated with Normal or Tenth Normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutralisation. 1 cc of Pyridin requires about 12.4 cc of Normal Hydrochloric Acid. 1 cc of Normal Volumetric Hydrochloric Acid Solution represents 0.07849 gramme of absolute Pyridin. The sample should not redden Phenolphthalein Solution, indicating the absence of Ammonia. It should have little or no action on Potassium Permanganate Solution, indicating the absence of readily oxidisable organic impurities, a 0.5 p c solution of Pyridin should give a crystalline precipitate, becoming almost insoluble when mixed with an equal volume of saturated Picric Acid Solution. It should be completely volatilised by heat without leaving any weighable residue.

PYROXYLINUM.**PYROXYLIN**

NO Syn —GOSYPIUM FULMINANS, FULMI COTON, LANA COLLODII, COLLOXYLINUM

FR, FULMI COTON, GR, KOTODIUMWOILE, ITAL, COLONT COLLODIO, SPAN, PIROXYLINA

A white, fibrous substance, having very much the appearance of ordinary cotton, but rapidly burning away with a flash when ignited. It requires to be kept in small quantities in a cool dry place, and away from lights.

Pyroxylin is Dinitrocellulose $C_6H_7(NO_2)_2O_5$, Gun Cotton is Trinitrocellulose $C_6H_5(NO_2)_3O_5$, and is not soluble in any mixture of Alcohol and Ether.

It sometimes decomposes on keeping, with disengagement of Nitrous fumes, and becomes insoluble.

The safest and best plan for its preservation is to moisten the dry material with an equal weight of Methylated Spirit, and preserve in a well stoppered jar, when required for use it is quickly and easily dried — *P J* '96, ii 110, *CD* '96, ii 207.

Pyroxylin is officially prepared by nitrating Cotton, 1 of Cotton being immersed in a mixture of 5 of Sulphuric Acid and 5 of Nitric Acid, the mixture being stirred during 3 minutes and the free acid separated by washing with Water till the washings no longer have an acid reaction. The Pyroxylin is drained and dried at a low temperature.

The proportions vary considerably in the different Pharmacopœias (see below).

Solubility —Readily soluble in a mixture of equal volumes of Ether and Alcohol (90 p c), also in Acetone.

Different samples of Pyroxylin vary considerably in the extent to which they are soluble.

The use of Acetone as a cheap and efficient solvent for Pyroxylin was suggested in the 16th edition of the *Companion*. A good Pyroxylin will dissolve readily to the extent of 10 p c. A formula corresponding with the *B P C* Collodion is Pyroxylin 1, Acetone 48, but this produces an inconveniently thin solution, and a preferable strength is Pyroxylin 3.5, Acetone 96.5 to make 100. The suggestion for its use has been followed by the *B P C* with certain additions to the formula to make it more closely resemble a well known proprietary article, it appears under the title **Collodium Acetinum** as follows — Pyroxylin 5, Oil of Cloves 2, Amyl Acetate 25, Benzol 20, Acetone 98 to produce 100. A more descriptive title for this would have been **Collodium Acetinum Compositum**, a simple solution in Acetone having been used as Acetone Collodion.

Foreign Pharmacopœias —Official in Belg (*Pyroxylinum*), Dutch, Ger and Jap, Nitric Acid 400, Sulphuric Acid 1000, Purified Cotton 55, Russ uses same formula, but Purified Cotton 50, Fr (*Fulmi coton*), Mex (*Piroxilina*), Cotton 1, Nitre 20, Sulphuric Acid 30, Port (*Algodao Polvora*), and Span (*Pyroxilina*), Cotton 1, Nitre 20, Pure Sulphuric Acid (sp gr 1.84) 30, Austr, Dan, Hung, Ital, Norw, Swed, Swiss and U S, no formula given.

Official Preparations —Used in the preparation of Collodium, and Collodium Vesicans. Of Collodium, Collodium Flexile.

Not Official —Collodium Stypticum, Hemostatic Collodion, Collodium Anodynum, Cellodion, Photoxylin.

Tests.—Pyroxylin should be readily and completely soluble in a mixture of equal volumes of Ether and Alcohol (90 p c), it should be neutral in reaction towards Litmus paper. It should also be soluble in Acetone. When ignited with free access of air it should leave no mineral residue.

Preparations.

COLLODIUM. COLLODION

Dissolve 1 of Pyroxylin in a mixture of Ether 36, and Alcohol (90 p c) 12, after a few days decant.

If the Pyroxylin be first soaked in the Alcohol, it quickly dissolves when Ether is added.—*P J '02, 1 188, C D '02, 1 269*

Mixes with Ether, but when mixed with Water or Alcohol (90 p c) the Pyroxylin is thrown out.

Official Preparation—Collodium Flexile

Official in Belg (Pyroxylum solutum), Pyroxylin 2, Alcohol 5, Ether 40, Castor Oil 3, Dan, Ger, Norw, Russ and Span (Collodium), Collodion Wool 4, Alcohol 12, Ether 84, Dutch, Collodion Wool 3, Alcohol 17, Ether 80, Fr and Mex, Collodion Wool 1, Alcohol 4, Ether 15, Ital (Collodio), Collodion Wool 1, Alcohol 4, Ether 12, Jap, Collodion Cotton 2, Alcohol 7, Ether 42, U S, Pyroxylin 4, Alcohol 25, Ether 75, Port, same as Collodio Elastico. All by weight except U S. Austr, Hung, Swed and Swiss, no formula given.

Tests.—Collodion has a sp gr of 0.770 to 0.780. Upon evaporation it leaves a thin elastic film, when evaporated to dryness and ignited with free access of air it leaves no weighable residue.

COLLODIUM FLEXILE. FLEXIBLE COLLODION. N O Syn — COLLODIUM ELASTICUM

Collodion, 12 fl oz, Canada Turpentine, $\frac{1}{2}$ oz (by weight), Castor Oil, $\frac{1}{4}$ oz (by weight).

Medicinal Properties.—Chiefly used for coating with a protective film, small clean cuts and abrasions, leech-bites, and fissure of nipple; it has been recommended as an application to erysipelatous surfaces and to burns, and to prevent the pitting of smallpox.

A large number of substances can be dissolved in Collodion to form medicated Colloids. See Acidum Salicylicum, Belladonna, Cantharis, Crotonis Oleum, Iodum, Iodoformum.

It does not contract or crack on drying.

Official in Austr, Dutch, Russ and Swed, Castor Oil 2, Collodion 98, Dan, Castor Oil 1, Collodion 99, Fr, Castor Oil 5, Collodion 95, Ger and Jap, Castor Oil 1, Turpentine 5, Collodion 94, Hung, Castor Oil 2, Collodion 100, Ital and Swiss, Castor Oil 3, Collodion 97, Mex, Castor Oil 9, Collodion 90, Norw, Glycerin 1, Collodion 99, Port, Pyroxylin 5, Castor Oil 5, Alcohol (90 p c) 20, Ether 70, Span, Castor Oil 10, Collodion 90, U S, Castor Oil 3, Canada Turpentine 5, Collodion 92. Not in Belg.

Tests.—Collodium Flexile has a sp gr of 0.790 to 0.800.

Not Official.

COLLODIUM STYPTICUM—Dissolve 44 grains of Benzoin in 1 fl. oz. of Absolute Alcohol, filter, and in the filtrate dissolve 1 oz of Tannic Acid, add Ether (sp gr 0.720), 4 fl oz, Pyroxylin, 44 grains, allow to stand 3 days, and decant.

B P C Formulary 1901, now incorporated in *B P C* as follows —

Benzoin 1 50, Pyroxylin 1 50, Tannic Acid 16, Alcohol 16, Purified Ether, *q s* to produce 100 — *B P C*

An adaptation of Dr. Richardson's Styptic Colloid

Official in U S, Tannic Acid 20, Alcohol 5, Ether 25, Collodion to 100

HÆMOSTATIC COLLODION (Dr. Prevost's) — Collodion 100, Carbolic Acid 10, Tannic Acid 5, Benzoic Acid 5, dissolve. Is applied by means of a camel hair pencil, or by soaking strips of linen in it

COLLODIUM ANODYNUM (Anodyne Collodion) — Aconitine, 1 grain, Veratrine, 6 grains, Æther Methylatus, 1 fl oz, Flexible Collodion, 1 fl oz

Collodium Anodynum *Syn* Anodyne Colloid — Aconitine 0 10, Veratrine 0 60, Flexible Collodion, *q s* to produce 100 — *B P C*

CELLOIDIN — A concentrated Collodion occurring in light, yellowish-brown, brittle strips. Is readily soluble in a mixture of Absolute Alcohol and Ether, and the solution is used for embedding histological specimens previous to cutting sections

A solution of Pyroxylin in Acetone is known under the name **Filmogen**

PHOTOXYLIN — A nitrated wood pulp prepared in St Petersburg. When made into Collodion it is stated to give a tougher film than Pyroxylin on evaporation. — *L* '87, 1 1253, *B M J* '88, 1 555

QUASSIÆ LIGNUM.

QUASSIA WOOD

FR, QUASSIA DE LA JAMAÏQUE, GLR, QUASSIAHOLZ, ITAL, QUASSIA, SPAN, LENO DI QUASSIA

The wood of the Trunk and Branches of *Picroëna excelsa*

Imported from Jamaica

It contains a bitter principle, **Quassin**, sparingly soluble in Water

Medicinal Properties — Possesses in a high degree the properties of the simple bitters, without astringency. For contra-indications, and other notes, see *Calumba*. Particularly valuable in dyspepsia due to the debility which succeeds acute disease, containing no Tannin, it is a compatible vehicle for Iron preparations. The infusion is also used as an anthelmintic enema in thread-worm

A few chips of Quassia or a weak infusion used in the morning bath is a protection against the annoying insects found in our cornfields. — *L* '84, 1 306. A strong infusion destroys fleas. — *L* '95, 1 1016

Official Preparations — Infusum Quassiae, Liquor Quassiae Concentratus, Tinctura Quassiae

Not Official — Extractum Quassiae, Fluidextractum Quassiae, Infusum Quassiae Concentratum

Foreign Pharmacopœias — Official in U S, same as Brit, Austr, Belg, Norw, Span, Swed and Swiss use *Quassia amara*, Dutch, Fr, Ger, Ital, Jap, Mex (*Quasia*), Port and Russ use both. Not in Hung or Dan. Fr has also Quassin

Descriptive Notes — In the *B P*, only the wood of the trunk and branches of *Picroëna excelsa*, Lindl (*Picroasma excelsa*, Planch) is official, but in the *P G* and *U S P* that of *Quassia amara*, L, is also official. The wood is met with in commerce in the form of splintery raspings or of coarse chips or transverse slices about an inch

in width and 1 to 4 in long and a line or more in thickness, but the official description refers only to impoised billets. These are usually 5 in or more in diameter. The wood should be nearly white, but is often yellowish or pale buff, it is easily cleft, but not hard. It has a bitter taste, but scarcely any odour. The medullary rays are usually 2 to 3 cells in width, *BP* (2 to 5 cells, *PG*, 3 to 5, *USP*), and 10 to 25 cells in height. In tangential section the cells of the medullary rays are seen to contain a series of single prisms of Calcium Oxalate.

The wood of *Quassia amara* (Suinam Quassia) has a deeper yellow colour, is harder and heavier, and the medullary rays are only 1 to 2 cells broad, and 5 to 20 cells high. The *PG* states that the wood is free from crystals of Calcium Oxalate, Vogl that there are no crystals in the wood, but sphaeraphides in the middle bark. Quassia that has been exhausted for the preparation of an agricultural insecticide has been offered in commerce, in this case the chips have dark lines of fungal hyphae present, and possess hardly any bitterness.

Tests.—Quassia contains about 3 p c of ash, and 4 p c should not be exceeded.

Preparations.

INFUSUM QUASSIÆ. INFUSION OF QUASSIA

Quassia Wood, finely rasped, 88 grains, Distilled Water, cold,
20 fl oz Macerate 15 minutes, and strain (about 1 in 100)

Dose — 1 to 1 fl oz = 14 2 to 28 4 c c

Foreign Pharmacopœias —Official in Fr (*Quassia Amara*), 1 in 200, Ital, 1 in 20 Spain (inct *Aleucosa de Quassia Amarga*), 1 in 100 Not in the others

A solid extract is official in the following Austri and Jap, first with cold Water, subsequently boiling, Fr (Quassia), Mex, Port and US use cold Water, Swiss uses boiling Water, Mex and US have also a fluid extract.

LIQUOR QUASSIÆ CONCENTRATUS. CONCENTRATED SOLUTION OF QUASSIA

2 of Quassia Wood, in No 40 powder, percolated with Alcohol
(20 p c), to produce 20 (1 in 10)

Dose.— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Tests.—Concentrated Quassia Solution has a sp gr of 0.975 to 0.980, contains about 0.5 p c w/v of total solids and about 18 p c w/v of Absolute Alcohol.

TINCTURA QUASSIÆ. TINCTURE OF QUASSIA

1 of Quassia Wood, rasped, macerated with 10 of Alcohol (45 p c)
(1 in 10)

Dose.— $\frac{1}{2}$ to 1 fl. drm = 1.8 to 3.6 cc

Foreign Pharm. Mex, Swiss ar Official in Belg, Dutch, Fr and Jap, 1 and 5, n the others All by weight except U.S

Tests.—Tincture of Quassia has a sp gr of 0.945 to 0.949; contains about 0.5 pc w/v of total solids and about 45 pc w/v of Absolute Alcohol. 0.016 pc w/v of Quassin has been suggested as a standard.

Not Official

EXTRACTUM QUASSIÆ —Quassia 16 is macerated with 8 of Water for 12 hours, exhausted by percolation, partly evaporated, filtered, and further evaporated until of a consistence for forming pills —*B P* 1885

This has been incorporated in the *B P C*

FLUIDEXTRACTUM QUASSIÆ —100 of Quassia, in No 40 powder, is exhausted with a mixture of Alcohol (95 p c) 30 and Water 60, the first 90 of percolate is reserved and the remainder evaporated to a soft extract, which is dissolved in the reserved portion, and enough menstruum added to make 100 —*U S P*

INFUSUM QUASSIÆ CONCENTRATUM —Quassia Wood, in No 20 powder, 7 5, Alcohol (90 p c), 20, Dilute Chloroform Water (1 in 100), *q s* to make 100 Prepare by repercolation

Dose — $\frac{1}{2}$ to 1 fl drim = 1 8 to 3 6 c c —*Farr* and *Wright*, *P J*, '06, 1 165 and '07, 1 623, *C D* '06, 1 252, *Y L P*, 1907, 249

This appears in the *B P C*, employing 5 of Quassia

Not Official

QUEBRACHO

The Bark of *Aspidosperma Quebracho*, Schlecht, obtained from the Argentine Republic and Brazil (*Quebracho blanco*)

Medicinal Properties —Was used rather extensively at one time for asthma and cardiac dyspnoea, but is now seldom prescribed

Foreign Pharmacopœias —Official in Austr, Mex, Span and Swiss
Not in the others

Tinctura Quebracho, 1 in 5 of Alcohol (60 p c), dose, $\frac{1}{2}$ to 1 fl drim = 1 8 to 3 6 c c

This has been incorporated in the *B P C*

Official in Mex, Span and Swiss

Fluid Extract (1 in 1) is official in Austr, and Extracto de Quebracho is official in Mex and Span

The following alkaloids and salts can be obtained Aspidospermine Cryst and Sulphate (Fraude), Aspidosamine and Hydrochloride (Hesse), Quebrachine Cryst and Hydrochloride (Hesse), dose, $\frac{1}{2}$ to 1½ grams, Quebrachamine and Sulphate (Hesse), Hypoquebrachine and Hydrochloride (Hesse)

Quebrachine is more active and more poisonous than Aspidospermine, it has greater antithermic properties —*L* '86, 1 804

Not Official

QUERCUS CORTEX

OAK BARK

The dried Bark of the small Branches and young Stems of *Quercus Robur*, L., collected in spring from trees growing in Britain

Medicinal Properties —A local astringent May be used topically in cases in which Tannic Acid is indicated, such as relaxed throat or tenderness of the gums, leucorrhœa, gonorrhœa, etc

Dose —Of the powder, 30 to 120 grains = 2 to 8 grammes Of a Decoction (1 to 16), 1 to 2 fl oz = 28 4 to 56 8 c c

Foreign Pharmacopœias —Official in Austr, Fr (Chêne), Ger, Hung, Mex (Encina), Norw, Port (Corvalho), Russ, Swiss, U S (*Quercus alba*)

QUILLAIAE CORTEX.

QUILLAYA BARK

B P Syn — PANAMA BARK *NO Syn* — SOAP BARK

FR, ÉCORCE DE PANAMA, GER, SEIFENRINDI, IAL, QUILLAJA,
SPAN, CORTEZA DE QUILLAYA

The inner part of the Bark of *Quillaja Saponaria*, Molina

Imported from Chili and Peru

Medicinal Properties—Has been strongly recommended as an expectorant, but its use requires caution, for it is a powerful irritant

The powder is excessively irritating to the passages

It has been found to possess properties allied to Senega, but it contains the two poisonous glucosides 'Quillaic Acid' and 'Sapotoxin' in much greater quantity than they exist in Senega

Prescribing Notes—*The Tincture is used to emulsify oils and fats, it requires from 1 to 3 of Tincture for 2 of Oil, depending on the character of the latter. Saponin is used for the same purpose*

Not Official—Saponin (Quillaic Acid)

Official Preparation—Tinctura Quillae Used in the preparation of Liquor Pice Carbonis

Foreign Pharmacopœias—Official in Austr, Dan (Quillaja), Fr (Écorce de Panama), Ger (Quillaja), Jap, Mex (Quillaya), Swiss and US (Quillaja) Not in the others US has a Fluid extract

Tests.—Quillaya Bark contains from 11 to 12 p c of ash

Preparation

TINCTURA QUILLAIAE. TINCTURE OF QUILLAYA

1 of Quillaya Bark, in No 20 powder, percolated with Alcohol (60 p c), to yield 20 (1 in 20)

Dose.— $\frac{1}{2}$ to 1 fl dm = 18 to 36 cc

Foreign Pharmacopœias—Official in Fr (Teinture de Panama), and Mex 1 in 5 (Alcohol 80 p c), Swiss, Quillaja 1, Alcohol 5, Water 5, US, boil 20 of Quillaja with 80 of Water for 15 minutes, strain and wash the residue with boiling Water, evaporate to 60, and when cold add 35 of Alcohol (95 p c), filter, and make up with Water to 100

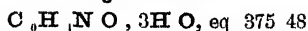
Tests.—Quillaya Bark Tincture has a sp gr of 0.920 to 0.925, it contains about 1.25 p c w/v of total solids and about 58 p c w/v of Absolute Alcohol

Not Official

SAPONIN (Quillain, Quillaic Acid)—A white, amorphous powder, soluble in Water and in Alcohol (90 p c), insoluble in Ether and in Chloroform

Not Official

QUININA



Fr, Hydramin de Quinini, Cir, Chinin, Lfai, Chinina,
Sian, Quinina

A white, soft, granular powder, slightly damp from adherent moisture, easily soluble in Ether or dilute mineral acids.

When separated from its solutions by shaking out with Ether or Chloroform and evaporating to dryness, it still retains a little Water, dried off with difficulty in a water bath, for determination purposes it should be heated to 120°C (250°F).

It should be kept in well stoppered glass bottles of a dark amber tint. When freshly precipitated from solutions of its salt it contains 3 molecules of Water of crystallisation as a Trihydrate.

Solubility Very sparingly in Water, 1 in 1 of Alcohol (90 p.c.), 1 in 3 of Chloroform, 1 in 4 of Ether.

Foreign Pharmacopœias - Official in Dutch, Fr, Hung, Mex, Port, Span, Swed and U.S.

Tests Quinine fuses when heated to 57°C (134.6°F) to a gummy looking mass. It loses 2 molecules of Water of crystallisation, equivalent to 9.2 p.c. when dried over Sulphuric Acid or heated at a temperature of 100°C (212°F), the remaining molecule of Water of crystallisation being driven off at 125°C (257°F), the total loss of Water being equivalent to 14.2 p.c. Quinine which has been rendered anhydrous by heating till constant in weight at a temperature of 125°C (257°F) melts at 175°C (347°F). The aqueous solution is alkaline in reaction towards moistened red Litmus paper, and is levogyrate. The alkaloid dissolves readily in diluted acids, its solution in diluted Sulphuric Acid exhibiting a strong fluorescence, the Hydrochloride and other haloid salts of Quinine exhibiting little fluorescence until excess of Sulphuric Acid is added. The fluorescence of solutions of the Sulphate is to a large extent destroyed by Hydrochloric Acid or by the presence of Chlorides. If just sufficient Sulphuric Acid be used to effect solution, and to this liquid be added 2 c.c. of Chlorine or Bromine Water, the subsequent addition of Ammonia Water produces an emerald green coloration. If the alkaloid be dissolved in diluted Alcohol and sufficient diluted Sulphuric Acid be added to convert it into a solution of Quinine Acid Sulphate and the liquid be then heated to boiling point, and a saturated Iodine Solution be added slowly and cautiously, the liquid on cooling deposits bronze green crystals of Quinine Iodo sulphate which are insoluble in cold Water. The acidified solution of the alkaloid is precipitated by Potassium Mercuric Iodide (Mayer's) Solution, and by Iodo potassium Iodide (Wagner's) Solution. When separated from its solutions by shaking out with Ether or Chloroform and evaporated to dryness it still retains a little Water, which is dried off with difficulty on a water bath. For determination purposes it should be heated to 120°C (250°F) before weighing. Quinine in the free state may be determined by titration with Tenth normal Volumetric Sulphuric Acid Solution, using Cochineal, Methyl Orange or Hematoxylin Solution as an indicator of neutrality. The behaviour of Quinine to these indicators of neutrality is somewhat anomalous, the point of neutrality when Cochineal or Hematoxylin is used as an indicator is reached when sufficient Acid has been added to convert the Quinine into the ordinary Quinine Sulphate ($\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2$) H_2SO_4 . That is to say that 1 c.c. of Tenth normal Volumetric Sulphuric Acid Solution is equivalent to 0.087848 gramme of Quinine Trihydrate or 0.032184 gramme of anhydrous Quinine. In the case of Methyl Orange the end reaction only occurs with the formation of the readily soluble Acid Sulphate ($\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2 \cdot \text{H}_2\text{SO}_4$), 1 c.c. of Tenth-normal Volumetric Hydrochloric Acid corresponding to 0.018774 gramme of Quinine Trihydrate or 0.016092 gramme of anhydrous Quinine. The use of Hydrochloric Acid to a large extent prevents the troublesome fluorescence yielded by the Sulphuric Acid Solution. Quinine may be distinguished from Cinchonine and Cinchonidine by the intensely red coloration, slowly changing to blue and finally to green, which is produced when 1 drop of Copper Sulphate

and a drop of Hydrogen Peroxide Solution is added to a gramme of the alkaloid in 20 c.c. of Water containing 1 c.c. Acid, Quinine and Quinidine both yield a reaction with this test. The absence of Cinchonine and Cinchonidine is shown by the fact that a solution of 1 gramme of the alkaloid in a slightly warm mixture of 6 c.c. of Absolute Alcohol and 3 c.c. of Ether remains clear on cooling. When treated with Sulphuric Acid it should not acquire more than a faintly yellowish colour, indicating the absence or limit of readily carbonisable organic impurities, nor should it produce a red colour on the addition of Nitric Acid, indicating the absence of Morphine. When heated with Potassium Hydroxide Solution it should not evolve an odour of Ammonia, nor should the issuing vapour have an alkaline reaction towards moistened red Litmus paper. When dried till constant in weight at 125°C (257°F), it should not lose more than 14.2 p.c. When ignited with free access of air it should leave no weighable residue.

Quinine should be free from the other Cinchona alkaloids, and when dissolved in Alcohol and carefully neutralised with Normal Volumetric Sulphuric Acid Solution, using Hamatoxylin Solution as an indicator of neutrality, and evaporated to dryness, the residue should respond to the official test for absence of Cinchonidine, Cinchonine, Quinidine, Cupieine and amorphous alkaloids given under Quinine Sulphas. The *U.S.P.* states that 2 grammes of Quinine, which have been previously dried at 50°C (122°F) for 2 hours in a porcelain dish, dissolved in 20 c.c. of Alcohol (94.9 p.c.) and neutralised exactly with Sulphuric Acid, using Hamatoxylin Solution as an indicator of neutrality, when evaporated to dryness on a water-bath yields a residue which answers the *U.S.P.* test for absence of other Cinchona alkaloids given under Quinine Sulphas. The alkaloid is not official in the German Pharmacopœia.

The official Salts of Quinine (Hydrochlorides and Sulphate) are given under separate headings.

INJECTIO QUININÆ HYPODERMICA—Quinine Hydrate, 76 grains; Lactic Acid, 27 minims, or a sufficiency, Distilled Water, a sufficiency, rub the Quinine with 6 fl. drms. of the Water, and add the Lactic Acid so as to dissolve the Quinine, and form a solution neutral or only faintly acid to Litmus paper, and make the measure up to 1 fl. oz. with Distilled Water.

More recently the Acid Hydrobiomide has been used for this purpose, see p. 986.

OLEATUM QUININÆ—Quinine, 1, Oleic Acid (by weight), 3, rub the Quinine with a small quantity of the Oleic Acid in a warmed mortar to form a smooth paste, add the remainder of the Oleic Acid, previously warmed, and stir frequently until the Quinine is dissolved.—*U.S.P.*

This has been in the *B.P.C.* under the title **Oleum Quininae** with the *syn.* **Oleum Quininae**.

QUININE ARSENATE ($\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2 \cdot \text{H}_3\text{AsO}_4 \cdot 2\text{H}_2\text{O}$, eq. 498.62)—Silky needles, sparingly soluble in cold Water, soluble in boiling Water. It may be prepared by the interaction of equivalent quantities of Quinine Hydrochloride and Mono-potassium Arsenate. It contains 64.5 p.c. of anhydrous Quinine, 28.3 p.c. of Arsenic Acid, and 7.2 p.c. of Water of crystallisation.

Dose— $\frac{1}{10}$ grain = 0.0065 gramme.

Tests—Quinine Arsenate dissolves sparingly in cold Water. The solution affords on the addition of a small quantity of Bromine Water, followed by a slight excess of Ammonia Solution an emerald-green coloration. The saturated aqueous solution affords a reddish-brown precipitate on the addition of Silver Ammonio-Nitrate Solution. 0.5 gramme of the salt when ignited with free access of air should leave no weighable residue.

QUININE ARSENATE BASIC ($\text{C}_{20}\text{H}_{24}\text{N}_2\text{O}_2 \cdot \text{H}_3\text{AsO}_4 \cdot 8\text{H}_2\text{O}$, eq. 927.74)—Colourless prismatic crystals, very soluble in cold Water, more readily soluble in hot Water. It contains 69.4 p.c. of anhydrous Quinine, 15.2 p.c. of Arsenic Acid and 15.4 p.c. of Water of crystallisation.

Tests—Basic Quinine Arsenate dissolves only sparingly in cold Water. The solution yields with Silver Ammonio-Nitrate Solution a reddish-brown precipitate, and when acidified with Hydrochloric Acid and warmed to about

80° C (176° F) it yields with Hydrogen Sulphide a yellow precipitate, soluble in Ammonium Carbonate Solution or in Sodium Hydroxide Solution. The saturated aqueous solution when treated with a small quantity of Bromine Water yields on the addition of Ammonia Solution in slight excess an emerald green coloration. 0.5 gramme of the salt when ignited with free access of air should leave no weighable residue.

QUININÆ CARBOLAS The crystalline salt contains 77 p.c. of anhydrous Quinine. For extemporaneous preparations, the alkoid may be used, and the best proportions are Quinine, 1, Carbolic Acid, 1, melt, and cool.

Dose 2 grains = 0.13 gramme for children.

Quininæ Sulphocarbolas—A yellowish white powder, prepared by the interaction of Quinine Sulphate and Barium Paraphenolsulphonate. **Dose**, 1 to 5 grains = 0.06 to 0.32 gramme.

QUININÆ CITRAS Crystallises in delicate needles.

Various formulas are given for this salt, QCl , Q_2Cl , $Q_2Cl \cdot 7H_2O$ but the commercial salt corresponds more closely with $(C_{20}H_{24}N_2O)_2H_3C_6H_5O_7 \cdot 3H_2O$, eq. 887.91, containing 72.5 p.c. of Quinine.

Solubility 1 in 1200 of Water, slightly in Chloroform.

QUININÆ CITRAS EFFERVESCENS—Contains 2 p.c. of Quinine Citrate in combination with Effervescent Sodium Citro-Tartrate, *BP*—*BP C*.

Official in Mex

QUININÆ ETHYL-CARBONAS (Euquinine, Euchinine)—Light, odourless, almost tasteless, silky, crystalline needles, sparingly soluble in Water, soluble in Alcohol, in Ether and in Chloroform. Produced by the action of Ethyl chlor carbonate on Quinine.

Antipyretic and analgesic. Recommended as a substitute for Quinine, owing to its tastelessness, and found useful in the hectic fever of tuberculosis, in whooping cough, influenza, and malaria.—*B M J E* '96, ii 104, '99, i 100, '01, ii 16, *B M J* '97, ii 1734, *L* '97, ii 728.

Dose—5 to 10 grains = 0.32 to 0.65 gramme.

Official in Jap and Swiss

Tests—Quinine Ethylcarbonas melts at about 95° C (203° F). It dissolves in a solution of Sulphuric or Nitric Acid, producing solutions having a strong green fluorescence. When treated with Sulphuric or Hydrochloric Acid and a little Chlorine Water it yields on the addition of Ammonia Solution in slight excess an emerald green coloration, when its solution in Sulphuric Acid is mixed with Solution of Iodine it does not yield crystals of the Iodo sulphate. When warmed with Sodium Hydroxide Solution, cooled, and sufficient Iodine Solution added to form a slight but distinct excess of the solution, again warmed it yields an odour of Iodoform. When dissolved in diluted Nitric Acid Solution it should yield no turbidity or precipitate with Hydrogen Sulphide Solution nor with Barium Chloride Solution. When heated with free access of air it should leave no weighable residue.

Under the name of **Aristochin**, a Di Quinine Carbonic Ester, in the form of an odourless, almost tasteless powder, has been introduced. As an analgesic $\frac{1}{2}$ gramme = 3½ grains, or more generally $\frac{1}{2}$ gramme = 7½ grains, given 3 times, 3, 2 and 1 hour before the pain is expected to begin. In doses of $\frac{1}{2}$ gramme = 7½ grains as an antipyretic in malaria.—*B M J E* '04, i 55.

QUININÆ ET FERRI CHLORIDUM—In brown scales or in a dark brown powder, very soluble in Water. Used as a hæmostatic.

Dose—5 to 15 grains = 0.32 to 1 gramme.

QUININÆ FLUORIDUM—A white, or whitish, amorphous powder.

Dose.—1 to 2 grains = 0.06 to 0.13 gramme.

QUININE FORMATES—There are two Formates of Quinine, the neutral Quinine Formate, prepared by dissolving the requisite amount of Quinine in solution of Formic Acid and allowing to crystallise, and the basic Quinine

Formate, prepared by neutralising Quinine with the calculated amount of Formic Acid

NEUTRAL QUININE FORMATE $C_{20}H_{24}N_2O_4 \cdot HCO_2$, eq 413.18 — White, shining needles, readily soluble in W. Very unstable, it contains 77.88 p.c. of alkaloid

Tests —Neutral Quinine Formate melts at $95^\circ C$ ($203^\circ F$). It loses Formic Acid at $50^\circ C$ ($122^\circ F$). It dissolves in Water and the aqueous solution yields when acidified with diluted Sulphuric Acid and treated with a small quantity of Chlorine or Bromine Water and an excess of Ammonia Solution an emerald-green coloration. The salt should leave no weighable residue when ignited with free access of air.

BASIC QUININE FORMATE $(C_{20}H_{24}N_2O_4 \cdot HCO_2)_2$, eq 867.51 — Forms white, silky needles, containing about 75 p.c. of Quinine. It is moderately soluble in Water, more so in boiling Water, readily soluble in Alcohol (90 p.c.) and in Chloroform, sparingly soluble in Ether, and insoluble in fixed Oils.

Tests —Basic Quinine Formate melts at about $109^\circ C$ ($228^\circ F$). The *B.P.C.* give the m.p. as $32^\circ C$, but this is apparently an error for $132^\circ C$. The m.p. given by Lacroix (*Jour Pharm Chem* [6] 22, 90) is $132^\circ C$ ($269.6^\circ F$), but this was subsequently corrected by him to $109^\circ C$ ($228.2^\circ F$). When dissolved in Water and acidified with diluted Sulphuric Acid the solution yields on the addition of a little Chlorine or Bromine Water, and subsequent addition of a slight excess of Ammonia Solution, an emerald-green coloration. Its aqueous solutions are strongly laevorotatory. The optical rotation is -144.2° . Lacroix originally gave the optical rotation as -141.1° , and this is the figure which has been adopted by the *B.P.C.*, it was altered by Lacroix to -144.2° . It should leave no weighable residue with free access of air.

QUININÆ GLYCEROPHOSPHAS —There are two Quinine Glycerophosphates, one basic and one neutral.

The basic salt $(C_{20}H_{24}N_2O_4)_2 \cdot C_3H_7O_3H_2PO_4 \cdot 5H_2O$, eq 903.89 is the one in general use. In slender, white, crystalline needles, slightly soluble in Water, 1 in 200 of Alcohol (90 p.c.).

Useful chiefly in neuralgia and in convalescence.

Dose —2 to 8 grains = 0.13 to 0.52 gramme

Official in F1 (Glycerophosphate Basique de Quinine)

Kineurine is stated to contain this salt.

Tests —Basic Quinine Glycerophosphate loses its Water of crystallisation, equivalent to 9.8 p.c., at $100^\circ C$ ($212^\circ F$), and is converted into an anhydrous salt. It melts at about $145^\circ C$ ($293^\circ F$). The aqueous solution affords with Potassium Hydroxide Solution a white precipitate soluble in Ether. The filtrate from the precipitate is evaporated to dryness and ignited with the addition of a little Potassium Nitrate, and Potassium Nitrate yields a residue, which, when dissolved in Water and acidified with Nitric Acid, affords with Ammonium Molybdate Solution a yellow precipitate soluble in Ammonia and reprecipitated as a white precipitate on the addition of Magnesium Solution. When dissolved in Water and acidified with Sulphuric Acid on the addition of a small quantity of Chlorine or Bromine Water and the subsequent addition of Ammonia Solution in slight excess, an emerald-green coloration. When shaken with Absolute Alcohol, filtered, and the alcoholic solution evaporated to dryness it should leave no weighable residue. The aqueous solution should afford no immediate precipitate with Ammonium Molybdate Solution.

QUININÆ HYDRIODIDUM $(C_{20}H_{24}N_2O_4 \cdot HI)$, eq 448.74 —The neutral salt has about the same solubility in Water as the Sulphate, and dissolves freely in Alcohol and Ether. It is generally found as a yellowish, amorphous powder.

SYRUPUS QUININÆ HYDRIODIDI. —Acid Hydriodide of Quinine, 4 scruples, Syrup, to 10 fl. oz. Triturate the Quinine in a mortar and add the Syrup gradually, stirring constantly to dissolve the salt. —*Pharm Form*

Syrupus Quininæ Hydriodidi *Syn* Syrup of Iodide of Quinine — Quinine Hydriodide, 2, Distilled Water, 2, Syrup of Citric Acid, *q s* to produce 100 — *B P C*

QUININÆ HYDRIODIDUM ACIDUM ($C_{20}H_{24}N_2O_2 \cdot 2HI \cdot 5H_2O$, eq 665.04) — Crystallises in large laminae of a fine yellow colour and is soluble 1 in 20 of Water

Both have been given in chronic rheumatism and tuberculosis

Dose — 1 to 5 grains = 0.06 to 0.32 gramme

QUININÆ HYDROBROMIDUM — Colourless, silky crystals, neutral or slightly alkaline

It should be kept in well stoppered glass bottles of a dark amber tint

It is given (*P J* (3) v 303) with H_2O , and soluble 1 in 5 *Fr Codex* with H_2O , soluble 1 in 41.5. Our stock (May 1893) corresponded with $C_{20}H_{24}N_2O_2 \cdot HBr \cdot H_2O$, containing 76.5 p.c. of Quinine, and soluble about 1 in 55 of Water, after drying at $125^\circ C$, its original moisture was again absorbed rapidly from the atmosphere. *U S* (1882) gave the formula with $2H_2O$, and solubility 1 in 16 of Water, *U S P* now gives it with H_2O , and soluble 1 in 40 of Water at $25^\circ C$ ($77^\circ F$)

The Hydrobromide is preferred (*Pr lxxiii* 682) for oral administration in malaria, and where rapid action is required, hypodermic or intravenous injection may be employed

Dose — 1 to 5 grains = 0.06 to 0.32 gramme

Official in Belg, Fr, Mex, Port, Russ, Span, Swed, Swiss and U S

Tests — Quinine Hydrobromidum loses its Water of crystallisation, equivalent to 4.25 p.c., when heated to $100^\circ C$ ($212^\circ F$), and at a higher temperature $152^\circ C$ ($305.6^\circ F$) it commences to fuse, forming a syrupy liquid. It dissolves in Water, forming a solution which is neutral or but faintly alkaline in reaction towards red Litmus paper. This solution when acidified with dilute Sulphuric Acid assumes a strong blue fluorescence, and when treated with a small quantity of Bromine Water and an excess of Ammonia Solution an emerald green colour is produced. The aqueous solution treated with Ammonia Solution affords a white precipitate soluble in an excess of the reagent, and the precipitate is also soluble in Ether. The addition of Potassium or Sodium Hydroxide Solution to an aqueous solution of the salt affords a white precipitate, and if this precipitate be removed by shaking with Ether the aqueous liquid when treated with a few drops of Chlorine Water assumes a yellowish or reddish colour, and when shaken with Chloroform the colour passes into the chloroformic solution. A portion of the aqueous liquid after separation of the Quinine when acidified with diluted Nitric Acid affords with Silver Nitrate, Solution a yellowish curdy precipitate insoluble in Nitric Acid, practically insoluble in Ammonia Solution, readily soluble in Potassium Cyanide Solution.

The more generally occurring impurities are excess of moisture, readily charred organic impurities, Sulphates, and other Cinchona alkaloids. The salt should not lose more than 4.25 p.c. of moisture when dried till constant in weight at $100^\circ C$ ($212^\circ F$). A solution of the salt in concentrated Sulphuric Acid should not be coloured more than a pale yellow, no red coloration should be produced on treating the salt with Nitric Acid. An aqueous solution of the salt should not assume more than a faint turbidity on the addition of Barium Chloride Solution.

It may be distinguished from Morphine by the Nitric Acid test described above, Morphine producing a red coloration with Nitric Acid. A confirmatory reaction for Morphine is to add 1 dgm. of Quinine Hydrobromide to 5 c.c. of a saturated Potassium Ferricyanide Solution, 25 c.c. of Water, 15 drops of Ferric Chloride T.S., and 5 c.c. of diluted Hydrochloric Acid, no blue coloration should be developed in 5 minutes. The absence of other Cinchona alkaloids may be assured by the *U S P* test given under Quininæ Sulphas. 3 grammes of the Hydrobromide which has been previously dried at $50^\circ C$ ($122^\circ F$) for 2 hours should be dissolved in 30 c.c. of hot Water, 1.5 gramme of crystallised Sodium Sulphate gradually added and the liquid evaporated to dryness on a water-bath, it then being examined by the test there described.

SYRUP QUININÆ HYDROBROMIDI—Quinine Hydrobromide, 80 grains, Dilute Hydrobromic Acid, 3 fl drms, Syrup of Orange (*B P* 1898), to make 10 fl oz. —1 *Ph I*

Quinine Acid Hydrobromide, 2, Syrup of Orange, *q s* to produce 100 — *B P C*

QUININÆ HYDROBROMIDUM ACIDUM eq
536 18) —Colourless crystals, containing 60 p c of Quinine

Solubility —1 in 6 of Water

3 grains dissolved in 20 minims of warm Distilled Water injected into the carefully asepticised upper arm, in the treatment of chronic malarial fever. 6 injections on alternate days are usually required in a serious case — *B M J* '99, ii 85, '02, i 201, 439, '03, i 848, *Y B P* '02, 203

Solutions of the **Acid Hydrobromide** and **Acid Hydrochloride** (which latter salt was made official in the British Pharmacopœia 1898) are put up in hermetically sealed glass capsules, and may be obtained in white or in dark amber tinted glass. Each c c contains 3 grains of the Acid Hydrobromide or $7\frac{1}{2}$ grains of the Acid Hydrochloride

Dose —1 to 5 grains = 0.06 to 0.32 gramme. Best administered hypodermically

Official in Fr and Mex, *Diomhydrate de Quinine Neutrie*

Tests —Quinine Acid Hydrobromide loses its Water of crystallisation, equivalent to 10.0 p c, when heated. It dissolves readily in Water, forming a clear solution which possesses an acid reaction towards blue Litmus paper. The aqueous solution yields on the addition of Ammonia Solution a white precipitate. When acidified with diluted Sulphuric Acid Solution and mixed with a small quantity of Bromine Water it yields on the addition of Ammonia Solution a slight excess, an emerald-green coloration. When acidified with diluted Nitric Acid Solution it yields with Silver Nitrate Solution a yellow curdy precipitate insoluble in Nitric Acid, insoluble in Ammonia Solution, readily soluble in Potassium Cyan. The salt loses 10.0 p c when dried till constant in weight. It should yield no turbidity on the addition of diluted Sulphuric Acid. When ignited with free access of air it should leave no weighable residue. It should be free from other Cinchona alkaloids when neutralised and examined by the tests described under Quinine Hydrobromidum, and using a correspondingly increased amount of Sodium Sulphate.

QUININÆ HYDROCHLORO-SULPHAS —Glistening, white, silky, crystalline needles, or as a white, or yellowish-white, amorphous powder. Soluble 1 in 2 of Water, 1 in 7 of Alcohol (90 p c). On account of its greater solubility in Water, it has been recommended for hypodermic use.

Dose —1 to 5 grains = 0.06 to 0.32 gramme

Official in Mex and Span, Mex has also solution for hypodermic injection, 1 in 2

QUININÆ HYPOPHOSPHIS ($C_{20}H_{24}N_2O_7$, H_4PO_4 , eq 387.40) —Generally supplied as an amorphous powder, but it can be obtained in light, colourless, prismatic crystals.

Solubility —1 in 250 of Water, 1 in 40 of Alcohol (90 p c)

Dose —1 to 5 grains = 0.06 to 0.32 gramme

QUININÆ IODO-HYDRIDIDUM —A reddish-brown, "
soluble in Water and in Alcohol, it is obtained by adding Iodo-Potassium Iodiac Solution to a solution of a Quinine salt. Has been employed in syphilitic diseases.

Dose —1 to 4 grains = 0.06 to 0.26 gramme

QUININÆ LACTAS ($C_{20}H_{24}N_2O_7$, $C_3H_5O_3$, eq 411.21) —Colourless prismatic needles or white crystalline powder; soluble about 1 in 6 of Water, but there is much doubt about its solubility.

Used chiefly by hypodermic injection in 10 p c solution

Official in Mex

QUININÆ PHOSPHAS—In light, white, acicular crystals. It is stated (*P J* (3) xiii 234) that the English made salt has the formula $3C_6H_7N O 2H_3PO_4 6H_2O$, and the German salt $2C_6H_7N O H_3PO_4 4H_2O$, the former containing 76 p c and the latter 79 p c of Quinine

Solubility—1 in 420 of Water, 1 in 110 of Alcohol (90 p c)

Dose—1 to 5 grains = 0.06 to 0.32 gramme

QUININÆ SALICYLAS ($C_{18}H_{14}N O C_7H_5O_3$), H_2O , eq 935.58—White, crystalline, silky needles, prepared by the interaction of Quinine Sulphate with Sodium Salicylate. It is practically anhydrous, and contains 70 p c of Quinine

It should be kept in well stoppered glass bottles of a dark amber tint and exposed as little as possible to the air. The *U S P* gives the formula of the salt with half a molecule of Water of crystallisation

Solubility—1 in 630 of Water, 1 in 24 of Alcohol (90 p c), 1 in 25 of Chloroform

In 1 drgm doses of the liquor every 3 or 4 hours for a day or two, combined with a spray for the post nasal space, is useful (*B M J* '05, ii 252, 1181) in preventing acute middle ear suppuration from becoming chronic

Dose—1 to 5 grains = 0.06 to 0.32 gramme

Given in capsules, cachets, or pills

Official in Mex, Russ, Span and U S

Tests—Quinine Salicylate when heated commences to melt at $183^{\circ} C$ ($361.4^{\circ} F$). It dissolves slightly in Water, forming a solution which possesses an alkaline reaction towards red Litmus paper, and which yields on the addition of Ferric Chloride T.S. a violet coloration. The saturated aqueous solution when mixed with a small quantity of Chlorine or Bromine Water yields on the addition of Ammoniac Solution in slight excess an emerald green coloration. Sulphuric Acid containing one fifth of its volume of Formaldehyde Solution yields a pink coloration. When heated till constant in weight at $100^{\circ} C$ ($212^{\circ} F$) the *U S P* requires that the salt should lose not more than 2.0 p c, indicating the absence of an excess of Water. When dissolved in Water acidified with a few drops of Nitric Acid and separated from the liberated Salicylic Acid, the filtrate should yield no marked turbidity on the addition of Silver Nitrate Solution, nor with Barium Chloride Solution, indicating the absence of more than slight traces of Chlorides and Sulphates. The alkaloid separated by mixing 2 grammes of the salt with 10 cc of Distilled Water adding Ammonia Solution in slight excess and extracting with 3 successive quantities each of 25 cc, 20 cc and 10 cc of Ether, evaporating the ethereal solution to dryness on a water bath, and dissolving the residue in Alcohol, when neutralised with Normal Volumetric Sulphuric Acid Solution, using Hematoxylin Solution as an indicator, shall leave when evaporated to dryness a residue which shall respond to the *U S P* tests for absence of other Cinchona alkaloids given under Quininæ Sulphas. If the aqueous alkaline liquid remaining after the extraction of the Quinine be acidified with diluted Sulphuric Acid and the liberated Salicylic Acid separated, carefully measured and dried it should possess the m.p., and answer the characteristic tests given under Acidum Salicylicum. When ignited with free access of air the salt should leave no weighable residue

QUININÆ SALICYLAS EFFERVESCENS—Can be obtained containing 1 and 5 grains in each drgm of Quinine Salicylate

Saloquinine (Quinine Ester of Salicylic Acid)—Colourless crystals, or a white amorphous powder, insoluble in Water, soluble in Alcohol (90 p c). Antipyretic, antiseptic, and analgesic. Has been recommended in typhoid fever and in neuralgia. Is also stated to possess antirheumatic properties.—*B M J* '02, i 782, *Y B P* '02, 204

This tasteless substitute for Quinine has been further recommended *P. lxxiii* 682) in 15 to 20 grain doses in malaria

Dose.—15 to 30 grains = 1 to 2 grammes

Rheumatine (Salicylquinine Salicylate) —Colourless crystalline needles, or as a white amorphous powder, soluble about 1 in 2000 of Water, 1 in 15 of Alcohol (90 p c) Antirheumatic Useful in acute articular rheumatism — *B M J* '02, i 782, *Y B P* '02, 204, *P J* '01, ii 645, *C D* '02, i 820

Dose —15 to 30 grains = 1 to 2 grammes

QUININÆ SULPHAS ACIDUS ($C_{20}H_{24}N_2O \cdot H_2SO_4 \cdot 7H_2O$, eq 544.34) — Translucent, colourless, or white, rhombic crystals It was originally called the **Neutral Quinine Sulphate**

It should be kept in well-stoppered glass bottles of a dark amber tint, as it has a tendency to effloresce on exposure to air

Solubility —1 in 10 of Water, 1 in 45 of Alcohol (90 p c)

Dose —2 to 12 grains = 0.13 to 0.78 gramme

A solution of 1 or 2 grains to the fl oz of Distilled Water applied to the eyes and nostrils for hay fever

2 grains twice daily as a prophylactic of influenza — *B M J* '02, i 940

5 grains injected into the subcutaneous tissue at the angle of the scapula repeated every 3 days in malaria Strong acids, especially Sulphuric, used to dissolve the Quinine salt may produce a local necrosis without agency of micro-organisms — *B M J* '02, i 1113

50 minims of a 1 in 5 solution successfully injected into each broad ligament in a case of prolapsus uteri — *B M J* '03, i 366

Foreign Pharmacopœias —Official in Austr and Hung (Chininum bisulphuricum), Dutch (Chininum bisulphas), Fr (Sulfate neutre de Quinine), Ital (Bisolfato di Chinina), Jap and US (Quininæ bisulphas), Mex (Sulfata de Quinina neutro), Span (Sulfato Quinico neutro)

Tests —Quinine Bisulphate when heated to $100^\circ C$ ($212^\circ F$) loses its Water of crystallisation, the loss being 23.2 p c It dissolves readily in Water, forming a solution which is acid in reaction towards Litmus paper, but which is neutral in reaction towards Methyl Orange Solution The solution exhibits a strong blue fluorescence, and when treated with a small quantity of Bromine Water it yields on the addition of Ammonia Solution in slight excess an emerald-green coloration It also yields on the addition of Barium Chloride a white precipitate insoluble in Hydrochloric Acid It should not lose more than 23.2 p c when dried at $100^\circ C$ ($212^\circ F$) The solution in Sulphuric Acid should not be of a deeper tint than a faint yellow, indicating the absence of readily charred organic impurities The absence of Cinchona alkaloids other than Quinine may be assured by dissolving 2 grammes of the salt which has been dried at $50^\circ C$ ($122^\circ F$) in 20 c c of Distilled Water, and after carefully neutralising the solution with diluted Sodium Hydroxide T S, evaporating to dryness on a water-bath, and examining the residue as directed under Quininæ Sulphas When ignited with free access of air it should leave no weighable residue

QUININÆ TANNAS —A yellowish-white, amorphous powder, sparingly soluble in Water, 1 in 3 of Alcohol (90 p c)

It should be kept in well-stoppered glass bottles of a dark amber tint and protected as far as possible from light It contains from 30 to 32 p c of anhydrous Quinine

Recommended because of its being tasteless

Dose.—1 to 10 grains = 0.065 to 0.65 gramme.

Official in Austr, Dan, Dutch, Ger, Hung, Jap, Mex., Norw., Port, Russ, Span and Swiss

Tests —Quinine Tannate when mixed with diluted Sulphuric Acid, a little Bromine Water and a slight excess of Ammonia Solution yields an emerald-

green coloration. The *P G* officially requires the salt to contain 30 p c of Quinine as gravimetrically determined by mixing 1 gramme of the Quinine Tannate with 4 c c of Water, adding Sodium Hydroxide Solution (15 p c) until strongly alkaline in reaction and extracting the mixture with 3 successive quantities each of 7 c c of Ether, evaporating the mixed ethereal solutions to dryness and drying the residue until constant in weight at 100° C (212° F). The Quinine obtained from the Tannate when exactly neutralised with diluted Sulphuric Acid should respond to the tests for other Cinchona alkaloids given under Quininae Sulphas. When shaken with diluted Nitric Acid and filtered, the filtrate should remain unaltered by the addition of Hydrogen Sulphide, Silver Nitrate Solution or Barium Chloride, indicating the absence of Copper and Lead, Chlorides and Sulphates. 0.2 of a gramme of the salt when ignited with free access of air should leave no weighable residue.

QUININÆ TARTRAS $(C_{20}H_{24}N_2O_6 \cdot C_{12}H_{10}O_6 \cdot H_2O, \text{eq } 810.48)$ —A white, crystalline powder.

Solubility—Very sparingly in Water (about 1 in 1000).

Quinine Sulphate, 90 grains, Tartaric Acid, 40 grains, Distilled Water, to measure 4 fl drms, has been used in India for hypodermic injection.

QUININÆ VALERIANAS $(C_{20}H_{24}N_2O_6 \cdot C_8H_{10}O_2, \text{eq } 423.15)$ —White, lustrous, pearly crystals, having an odour of Valerianic Acid. Can be prepared by decomposing Quinine Hydrochloride with Sodium Valerianate.

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from the light.

Solubility—1 in 120 of cold Water, 1 in 2 of Alcohol (90 p c), 1 in 14 of Ether.

Dose—1 to 3 grains = 0.06 to 0.2 gramme.

Official in Fr, Ital, Mex, Port, Spain and Swed.

Tests—Quinine Valerianate melts when heated to about 90° C (194° F), at 100° C (212° F) it loses Valerianic Acid pretty rapidly. It dissolves slightly in cold Water, the solution being neutral or faintly acid in reaction towards Litmus paper. The aqueous solution affords with Ammonia Solution a white precipitate soluble in excess of the reagent. It affords when treated with a small quantity of Bromine Water and an excess of Ammonia Solution an emerald green coloration. When acidified with diluted Sulphuric Acid Solution it evolves a characteristic odour of Valerianic Acid, the solution exhibiting a blue fluorescence. The alkaloid extracted from the salt by treatment with Ether in alkaline solution when carefully neutralised with Sulphuric Acid, using Hæmatoxylin Solution as an indicator of neutrality should respond to the test for freedom from other Cinchona alkaloids given under Quininae Sulphas. The salt should not yield more than a faint yellow tint when mixed with concentrated Sulphuric Acid, indicating the absence of readily charred organic impurities. An aqueous solution of the salt should not be rendered distinctly turbid by Barium Chloride Solution. When ignited with free access of air it should burn without leaving a weighable residue.

Quinine Camphorate, a white powder insoluble in Water, soluble in Alcohol (90 p c), dose, 1 to 10 grains = 0.06 to 0.65 gramme, **Quinine Bihydrochloro carbamide**, prismatic crystals soluble in Water, dose, 5 to 15 grains = 0.32 to 1 gramme, chiefly used hypodermically, **Quinine Saccharinate** (Basic), crystalline needles insoluble in Water, **Quinine Sulphocarbolate**, a yellowish-white powder soluble in Alcohol, dose, 1 to 5 grains = 0.06 to 0.32 gramme, **Quinine Sulphocresotate**, yellow scales soluble in Water, dose, 1 to 5 grains = 0.06 to 0.32 gramme, and **Quinine Vanadate** are salts of Quinine which have received some attention in medical literature.

SYPHUS QUININÆ DIKINATIS—Introduced by Dr Donovan of Dublin.

1 fl drms contains 2 grains of Quinine Dikinate.

Dose— $\frac{1}{2}$ to 1 fl drms = 1.8 to 3.6 c c.

WARBURG'S TINCTURE FOR MALARIAL FEVER—The formula for this is given in the *MT* '75, ii 540, with some interesting cases by Professor

Maclea, C B, Aloes Socotrinae 4, Rad Rhei 4, Sem Angelicae 4, Conf Damocratis 4, Rad Heleni 2, Cicut Sativi 2, Sem Foeniculi 2, Cretae Preparatae 2, Rad Gentianae 1, Rad Zedoariae 1, Pip Cubebe 1, Myrrh Elect 1, Camphora 1, Bolet Larici 1 Digest with 500 of Proof Spirit on a water-bath for 12 hours, express, and add Quinine Sulphatis 10 Continue heating on a water-bath till all the Quinine Sulphate is dissolved, filter when cold

Warburg's Tincture is without its equal in persistent and protracted agues — *T G '94, 842*

A somewhat similar preparation was included in the *B P C Formulary* 1901 under the title *Tinctura Antiperiodica* as follows —

Tinctura Antiperiodica. *Syn* Warburg's Tincture — Socotrine Aloes, bruised, 240 grains, Rhubarb, bruised, 80 grains, Angelica Fiut, bruised, 80 grains, Elecampare Root, bruised, 40 grains, Saffron, 40 grains, Fennel, bruised, 40 grains, Prepared Chalk, 40 grains, Gentian, bruised, 20 grains, Zedoary Root, bruised, 20 grains, Cubebs, bruised, 20 grains, Myrrh, elect and bruised, 20 grains, White Agaric, in powder, 20 grains, Opium, in powder, 2½ grains, Black Pepper, bruised, 4 grains, Cinnamon, bruised, 8 grains, (Ginger, bruised, 8 grains, Alcohol (60 per cent), a sufficient quantity

Macerate for 7 days in 1 pint of the Alcohol, press and filter Dissolve in the product — Quinine Sulphate, 175 grains, Camphor, 20 grains After 3 days filter, and add sufficient of the Alcohol to make 1 pint

Dose — 1 to 4 fl drms

This has been incorporated in the *B P C*

QUINETUM — The mixed alkaloids from the E I Red Cinchona Bark The Sulphate resembles Quinine Sulphate

Solubility — Sparingly in Water, 1 in 90 of Alcohol (90 p c)

Dose — Of the Sulphate 1 to 10 grains = 0.06 to 0.65 grammes

QUINIDINÆ SULPHAS ($C_{20}H_{24}N_2O_4$) \cdot $H_2SO_4\cdot 2H_2O$, eq 776.78 — White, silky crystals It should be kept in well-stoppered bottles

Solubility — 1 in 200 of Water, 1 in 24 of Alcohol (90 p c), about 1 in 100 of Glycerin

Dose — 10 to 20 grains = 0.65 to 1.3 grammes

QUINOIDIN *Syn* CHINOIDIN — A mixture of Alkaloids, mostly amorphous, obtained as a by-product in the manufacture of the crystallisable alkaloids from Cinchona A brownish-black mass with alkaline reaction On ignition should not leave more than 0.7 p c of ash

Official in Spain

QUINOLINE Chinoline C_9H_7N , eq 128.18 — It is formed by the distillation of Quinine or Cinchonine with aqueous Potassium Hydroxide, or synthetically from Aniline and Nitrobenzene It is a colourless, mobile liquid, having a faint aromatic odour and a peculiar penetrating taste, sparingly soluble in Water, miscible with Alcohol, Ether and Carbon Bisulphide It should be preserved in well-stoppered bottles of an amber tint

Dose — 5 to 15 grains = 0.32 to 1 gramme

CHINOLINE PERIODIDE — Chinoline may be produced synthetically from Aniline and Nitrobenzene, or by the distillation of Quinine The above Iodide is one of the series of Iodides introduced by Squibb at the suggestion of Dr. Mortimer Granville and employed in the treatment of gout The Chloride used in this preparation is not of synthetic production — An Iodide made with Chinoline, prepared from Cinchonine, is known as **Cincho-quinoline Periodide** (*Squire*)

LORETIN (Methiod-oxaloxycarbonyl-aminophenyl acid) — A pale yellowish powder, odourless and non-poisonous Introduced as a substitute for Iodoform Used as a dusting powder and the correct Quinine *B P C '98, 11 91; M A '95, 34, L '94, 11 31 '95, 11 153, M P '94, 11 20*

CHINOSOL (Quinosol Potassium Oxychinoline Sulphonate)—A bright lemon yellow powder with a faint odour, soluble in Water

A powerful antiseptic, disinfectant and deodorant. Action more marked as a lotion than as a powder. When used as a powder should be diluted. Solutions for disinfection of instruments should not be too concentrated. Drug possesses toxic properties. If used in too concentrated a form subcutaneously it will produce local irritation and swelling. Is not rapidly absorbed by the unbroken skin.—*B M J* '98, i 91

In doses of 1 to 5 grains internally and as a local application it has given good results in leprosy.—*P J* '99, ii 135

5 grains 3 times a day after food in the treatment of over one hundred cases of pulmonary phthisis. In almost every case improvement in the patient's general condition followed.—*L* '99, ii 90, 181, 238

1 to 2 p.c. solution has power of arresting hæmorrhage.—*B M J E* '01, ii 60

Official in Russ

Cruin (Quinoline Bismuth Sulphocyanide)—A yellowish red powder, in soluble in Water and Alcohol. Recommended in $\frac{1}{2}$ p.c. solution as an injection in gonorrhœa.—1 of Cruin rubbed up with Glycerin and Water, of each 5, and made up with Water to 200.—*B M J E* '02, i 32, *C D* '02, i 643, *P J* '00, i 615, '00, ii 486, '02, i 442

Vioform (Iodochloroxychinoline, Iodochloroxyquinoline)—An almost odourless, non-toxic powder, insoluble in Water. Antiseptic and germicide. Introduced as a substitute for Iodoform. It has been found useful in operations upon tubercular joints. Most conveniently used as an emulsion.—Vioform 50, Glycerin 200, Sterilised Water 200, Alcohol 100.—*B M J E* '03, i 31, *P J* '00, ii 470, 700, '02, i 513, *B M J E* '07, i 100

Diaphthol (Quinaseptol) and **Diaphtherin** (Oxychinaseptol) have also been used as antiseptics

QUININÆ HYDROCHLORIDUM.

QUININE HYDROCHLORIDE

HYDROCHLORATE OF QUININE.—*B P* '85

$C_{20}H_{24}N_2O_2.HCl, 2H_2O$, eq 398.79

FR, CHLORHYDRATE BASIQUE DE QUININE, GER, CHININHYDROCHLORID, ITAL, CLORIDRATO DI CHININA, SPAN, CLORURO QUINICO

White, odourless, silky, needle-shaped crystals, possessing a very bitter taste, and which have a tendency to lose Water in warm air. It is officially described as the Hydrochloride of an alkaloid obtained from the Bark of various species of *Cinchona* and *Remaja*. The *U S P* describes it as the Hydrochloride of the alkaloid Quinine.

It should be kept in well stoppered glass bottles of a dark amber tint.

The salt contains theoretically 81.7 p.c. of anhydrous Quinine, 9.2 p.c. of Hydrochloric Acid and 9.1 p.c. of Water of crystallisation.

Solubility—1 in 37 of Water, 1 in 1 of boiling Water, 1 in 1 of Alcohol (90 p.c.). The anhydrous salt is very soluble in Chloroform.

Medicinal Properties—Same as Quinine Sulphate.

This salt is preferred for the prevention of ague for the following reasons: (1) it is more readily soluble and very easily absorbed, (2) it is less irritating to the gastric mucous membrane, (3) it contains relatively a greater proportion of Quinine, (4) it is the chief soluble salt of Quinine, and is almost universally used in the malarial districts of Italy.—*B M J E* '03, ii 12

Has been shown (*B M J* '04, ii 1543) to maintain the heart's action during

an operation when administered in doses of a few grains, 24 hours before the operation

Two cases of idiosyncrasy to Quinine are noted (*Pr Lxxiii* 682) in which the Sulphate produced alarming symptoms, whilst the Hydrochloride was well borne

Topical use in leucorrhœa, 2 to 3 grains as a pessary —*L* '99, 1 26, 192

As a styptic and antiseptic agent Recommended for parenchymatous hæmorrhages —*L* '01, 11 1541

Inoperable cancer of the uterus, successfully treated by endovenous injection of 4 to 8 grains —*B M J E* '03, 1 26

Dose.—1 to 10 grains = 0.06 to 0.65 gramme

Official Preparations —Tinctura Quininae and Vinum Quininae

Not Official —Pessus Quininae, Solutio de Quinine pour Injection Hypodermique

Foreign Pharmacopœias —Official in Austr., Ger., Hung., Jap., Russ. and Swiss (*Chininum Hydrochloricum*), Belg. (*Chlorhydrate de Quinine*), Dan., Norw., Swed. (*Chloridum Chinicum*), Dutch (*Hydrochloras Chinini*), Fr. (*Chlorhydrate Basique de Quinine*), Ital. (*Cloridrato di Chinina*), Mex. (*Clorhidrato de Quinina basico*), Port. (*Chlorhydrate de Quinine*), Span. (*Cloruro Quinico*), U.S. (*Quinine Hydrochloridum*)

Tests.—Quinine Hydrochloride when heated to a temperature of 100° C (212° F) loses 9 p.c. of Water equivalent to 2 molecules of Water of crystallisation. The *USP* states it loses its Water of crystallisation at a temperature of 120° C (248° F), and that at about 156° C (312.8° F) it commences to melt, but that it is not fully melted until a temperature of 190° C (374° F) is reached. It dissolves fairly readily in Water, forming a solution which is neutral to Litmus paper or at the most but faintly alkaline in reaction towards red Litmus paper. On the addition of Sulphuric Acid the aqueous solution assumes a strong bluish-green fluorescence, but the solution of the Hydrochloride itself is not fluorescent. The alkaloid extracted from a solution of the Hydrochloride should answer the tests distinctive of Quinine given under that substance. The aqueous solution when acidified with Nitric Acid yields with Silver Nitrate Solution a white curdy precipitate, which, when washed, dissolves readily and completely in Ammonia Solution. The percentage of Quinine may be determined by the direct titration of a solution of the Hydrochloride with Tenth-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality, 1 c.c. of the Tenth-normal Volumetric Solution being equivalent to 0.039379 gramme of the crystallised Hydrochloride. The alkaloidal content may be gravimetrically determined by dissolving the salt in Water, adding sufficient Potassium or Sodium Hydroxide Solution to render the liquid distinctly alkaline, and shaking out with Ether-Chloroform Solution.

The more generally occurring impurities are excess of moisture, Sulphates, readily charred organic impurities, Cinchona alkaloids other than Quinine, and mineral impurities. The *BP*, the *USP* and the *PG* require that the salt shall not lose more than 9 p.c. of weight when dried at a temperature of 100° C (212° F). An aqueous solution of the salt should not be rendered more than slightly turbid on the addition of Barium Chloride Solution, indicating the limit of

Sulphate It should produce no coloration on the addition of Sulphuric Acid, indicating the absence of readily charred organic impurities. The *BP* requires that the salt should yield only the slightest characteristic reactions with the tests for Sulphates, and that when the Hydrochloride is converted into the Sulphate by mixing it with an equal weight of Sodium Sulphate and dissolving the mixture in 10 times its weight of hot Water, allowing the mixture to stand at 15° C (60° F), it should answer the tests described under Quinine Sulphas. The *USP* requires that a weighed quantity of 3 grammes of the salt, which has been previously dried for 2 hours at a temperature of 50° C (122° F), when dissolved in 30 c.c. of hot Distilled Water, mixed with 1.5 grammes of crystallised Sodium Sulphate, gradually and with constant stirring, and the liquid evaporated to dryness on a water-bath, the residue when dissolved in 30 c.c. of Water should respond to the *USP* test for absence of an excessive amount of Cinchona alkaloids other than Quinine. The *PG* dissolves a weighed quantity of 2 grammes of the Hydrochloride in a warmed mortar in 20 c.c. of Water at a temperature of 60° C (140° F). To the solution is added 1 gramme of powdered uneffloresced Sodium Sulphate, and the mixture thoroughly incorporated. It is allowed to stand when cold for half an hour at a temperature of 15° C, it is then pressed through a dry piece of calico of about 100 cm. square, and the expressed fluid filtered through a piece of the best filter paper. A measured quantity of 5 c.c. of this filtrate is brought to a temperature of 15° C (59° F), and mixed with Ammonia Solution at a temperature of 15° C (59° F) until the precipitate, which at first separates out, again dissolves to a clear solution, not more than 4 c.c. of Ammonia Solution shall be required. 1 gramme of the salt when ignited with free access of air should leave no weighable residue, indicating the absence of mineral impurities. Quinine may be distinguished from Morphine by the Nitric Acid colour test, the salt should dissolve in Nitric Acid without the production of a red colour. The *PG* states that 0.5 of a gramme of the salt mixed with 10 drops of Sulphuric Acid and 1 drop of Nitric Acid shall not yield a reddish-yellow coloration. The *USP* includes an additional test for differentiation from Morphine, it directs that 0.1 of a gramme of the salt added to 5 c.c. of a saturated Potassium Ferricyanide Solution, 25 c.c. of Water, 15 drops of Ferric Chloride T.S. and 5 drops of diluted Hydrochloric Acid Solution should not produce a blue coloration after being well shaken and allowed to stand for 5 minutes.

Residue—When dried at 212° F (100° C), 1 gramme of the salt should not lose more than 0.09 gramme in weight, *BP*, *PG* and *USP*. After ignition it should leave no residue, *PG* and *USP*.

Litmus—Its aqueous solution is neutral, *PG*, or faintly alkaline, *USP*.

Barium Nitrate or Chloride—An aqueous solution of the salt should not be rendered more than faintly turbid by T.S. of Barium Chloride, *USP*, by T.S. of Barium Nitrate, *PG*.

Sulphuric Acid—No turbidity at all should be produced in an aqueous solution (1-50) of the salt by diluted Sulphuric Acid, *PG*. The salt should not

yield any colour with Sulphuric Acid, *USP* 0.05 gramme of the salt mixed with 10 drops of Sulphuric Acid and 1 drop of Nitrate Acid, should not assume a reddish yellow colour, *PC*

Preparations

TINCTURA QUININÆ. TINCTURE OF QUININE

Quinine Hydrochloride, 175 grains, Tincture of Orange, 20 fl oz
(about 1 grain in 55 minims)

Dose — $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 cc

Tests — Tincture of Quinine has a sp gr of 0.880 to 0.890, it contains about 3.5 p.c w/v of total solids and about 74 p.c w/v of Absolute Alcohol

VINUM QUININÆ QUININE WINE

Quinine Hydrochloride, 20 grains, Orange Wine, 20 fl oz

Dose. — $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc

Now made with Quinine Hydrochloride instead of Sulphate

Tests — Quinine Wine has a sp gr of 1.044 to 1.095, it contains about 16 p.c w/v of total solids and from 10 to 12 p.c w/v of Absolute Alcohol. It should yield 0.187 p.c w/v of anhydrous Quinine. 1 fl oz of the wine when made alkaline with Sodium Hydroxide Solution and shaken with Ether, the aqueous alkaline layer separated, and after acidification shaken with a further quantity of Ether, the ethereal solution when mixed with a little Water, a drop or two of Ferric Chloride Solution added and the mixture well shaken should yield no violet coloration, indicating the absence of Salicylic Acid

Not Official

PESSUS QUININÆ — 3 to 5 grains of Quinine Hydrochloride. A valuable remedy for leucorrhœa. — *Martindale*

This has been incorporated in the *BPC*

SOLUTÉ DE QUININE (CHLORHYDRATE BASIQUE) POUR INJECTION HYPODERMIQUE — Basic Hydrochloride of Quinine, 3 grammes, Antipyrine, 2 grammes, Distilled Water, boiled and cooled, *qs* to obtain 10 cc of solution. — *Fr*

QUININÆ HYDROCHLORIDUM ACIDUM.

ACID QUININE HYDROCHLORIDE

$C_{20}H_{24}N_2O_2 \cdot 2HCl, 3H_2O$, eq 447.86

Fr, CHLORHYDRATE NEUTRE DE QUININE, *GER*, SAURES CHININHYDROCHLORID, *ITAL*, BICHLORIDRATO DI CHININA

Small, colourless, glistening crystals, or as a white, odourless, crystalline powder, possessing a very bitter taste. It is officially described as the Acid Hydrochloride of an alkaloid obtained from the Bark of various species of *Cinchona* and *Remijna*, but would have been better described as the Acid Hydrochloride of the alkaloid Quinine

It should be kept in well-stoppered glass bottles of a dark amber tint and exposed as little as possible to the air. The official formula for the salt shows 3 molecules of Water of crystallisation but the majority of commercial specimens

contain practically no Water of crystallisation. Howard states that the salt is anhydrous if dried at 100°C (212°F), but that the Pharmacopœia formula is correct for the crystalline salt formed at a lower temperature. The *Fr Codea* (1908) gives the formula with $2\frac{1}{2}$ molecules of Water of crystallisation, and states that from Absolute Ethylic Alcohol it forms acicular crystals containing 1 molecule of Alcohol of crystallisation which it loses readily. The dried salt when exposed to the air reabsorbs moisture, equivalent to $2\frac{1}{2}$ molecules of Water of crystallisation.

Solubility—2 in $1\frac{1}{2}$ of Water and measures 3, 1 in 5 of Alcohol (90 p c), 1 in 7 of Chloroform. Insoluble in Ether.

Medicinal Properties—Same as Quinine Sulphate and Hydrochloride. It is frequently employed by hypodermic injection. See notes on the Acid Hydrobromide.

Intramuscular injections in malaria.—*J*, '02, 1 1379

5 to 10 grains twice a day for six weeks injected into the gluteal muscles in the treatment of ague.—*B M J* '02, 11 1767

15 grains twice daily given on an empty stomach in the treatment of typhoid fever in the tropics.—*B M J E* '02, 1 80

In doses of 2 to 3 grains hypodermically in the treatment of blackwater fever.—*B M J* '02, 1 1334, *P J* '02, 11 249

1 to 2 grains injected into the subcutaneous tissue over the splenic area on 3 or 4 successive mornings in the treatment of malaria.—*B M J* '03, 1 848

Quinine Bihydrochloride is now exclusively recommended (*B M J* '06, 1 1398) in amœbic abscess of the liver. Two solutions are prepared and sterilised before the operation, each containing 30 grains of this salt, but in one this amount is dissolved in 2 oz of Water and in the other in 4 oz, the former being used if the abscess contains less than 10 oz of pus and the latter if it is larger. In this way the dose of the salt is limited to 30 grains.

Dose—1 to 10 grains = 0.06 to 0.65 gramme

Foreign Pharmacopœias—Official in *Fr* (Chlorhydrate neutre de Quinine), *Ital* and *Mex*

Tests—Quinine Bihydrochloride is officially required to lose not more than 12 p c, equivalent to practically 3 molecules of Water of crystallisation at a temperature of 100°C (212°F). It dissolves readily in Water, forming a clear solution which possesses a strong acid reaction towards Litmus, and which yields, on the addition of Potassium or Sodium Hydroxide Solution, a white precipitate, if this precipitate be separated and carefully washed it answers the tests distinctive of Quinine given under that heading. An aqueous solution of the salt, when acidified with diluted Nitric Acid, yields, on the addition of Silver Nitrate solution, a white curdy precipitate, insoluble in Nitric Acid, and which, when separated and washed, dissolves readily and completely in Ammonia Solution. The total percentage of Hydrochloric Acid present may readily be determined by titration with Tenth-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. In conducting the titration, sufficient neutral Ether may be added to hold the liberated alkaloid in solution, the end reaction not being then masked by the precipitate. 1 c c of Tenth normal Volumetric Alkali Solution is equivalent to 0.003619 gramme of Hydrochloric Acid and to 0.044786 gramme of crystallised Acid Quinine Hydrochloride of the official formula, or to 0.039422 gramme of the anhydrous Hydrochloride. 1 gramme of the salt, when dissolved in 20 c c of Water,

yield any colour with Sulphuric Acid, *U S P* 0.05 gramme of the salt mixed with 10 drops of Sulphuric Acid and 1 drop of Nitrate Acid, should not assume a reddish-yellow colour, *P G*

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FR, CHLORHYDRATE NEUTRE DE QUININE, GER, SAURES CHININHYDRO-
CHLORID, ITAL, BICHLORIDRATO DI CHININA

Small, colourless, glistening crystals, or as a white, odourless, crystalline powder, possessing a very bitter taste. It is officially described as the Acid Hydrochloride of an alkaloid obtained from the Bark of various species of *Cinchona* and *Remyria*, but would have been better described as the Acid Hydrochloride of the alkaloid Quinine.

It should be kept in well-stoppered glass bottles of a dark amber tint and exposed as little as possible to the air. The official formula for the salt shows 3 molecules of Water of crystallisation, but the majority of commercial specimens

contain practically no Water of crystallisation. Howard states that the salt is anhydrous if dried at 100°C (212°F) but that the Pharmacopœia formula is correct for the crystalline salt formed at a lower temperature. The *B. Codex* (1908) gives the formula with $2\frac{1}{2}$ molecules of Water of crystallisation and states that from Absolute Ethylic Alcohol it forms acicular crystals containing 1 molecule of Alcohol of crystallisation which it loses readily. The dried salt when exposed to the air reabsorbs moisture, equivalent to $2\frac{1}{2}$ molecules of Water of crystallisation.

Solubility—2 in 11 of Water and measures 3, 1 in 5 of Alcohol (90 p.c.), 1 in 7 of Chloroform. Insoluble in Ether.

Medicinal Properties—Same as Quinine Sulphate and Hydrochloride. It is frequently employed by hypodermic injection. See notes on the Acid Hydrobromide.

Intramuscular injections in malaria.—*J.* '02, i, 1379

5 to 10 grains twice a day for six weeks injected into the gluteal muscles in the treatment of ague.—*B. M. J.* '02, ii, 1767

15 grains twice daily, given on an empty stomach in the treatment of typhoid fever in the tropics.—*B. M. J.* '02, i, 80

In doses of 2 to 3 grains hypodermically in the treatment of blackwater fever.—*B. M. J.* '02, i, 1334 *P. J.* '02, ii, 249

1 to 2 grains injected into the subcutaneous tissue over the splenic vein on 3 or 4 successive mornings in the treatment of malaria.—*B. M. J.* '03, i, 848

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Dose—1 to 10 grains = 0.06 to 0.65 gramme

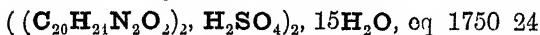
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Tests—Quinine Bihydrochloride is officially required to lose not more than 12 p.c., equivalent to practically 3 molecules of Water of crystallisation at a temperature of 100°C (212°F). It dissolves readily in Water, forming a clear solution which possesses a strong acid reaction towards Litmus, and which yields, on the addition of Potassium or Sodium Hydroxide Solution, a white precipitate, if this precipitate be separated and carefully washed it answers the tests distinctive of Quinine given under that heading. An aqueous solution of the salt, when acidified with diluted Nitric Acid, yields, on the addition of Silver Nitrate solution, a white curdy precipitate, insoluble in Nitric Acid, and which, when separated and washed, dissolves readily and completely in Ammonia Solution. The total percentage of Hydrochloric Acid present may readily be determined by titration with Tenth-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. In conducting the titration, sufficient neutral Ether may be added to hold the liberated alkaloid in solution, the end reaction not being then masked by the precipitate. 1 c.c. of Tenth-normal Volumetric Alkali Solution is equivalent to 0.003619 gramme of Hydrochloric Acid and to 0.044786 gramme of crystallised Acid Quinine Hydrochloride of the official formula, or to 0.039422 gramme of the anhydrous Hydrochloride. 1 gramme of the salt, when dissolved in 20 c.c. of Water,

is officially stated to require not more than 2.5 c.c. of Volumetric Sodium Hydroxide Solution for its complete neutralisation. Unfortunately the *B.P.* has omitted to mention to what indicator of neutrality, if Phenolphthalein Solution be used as an indicator, considerably more than 2.5 c.c. of Volumetric Solution will be required, as indicated above the whole of the acid radicle is determined by this indicator. With Litmus and Methyl Orange Solution the results are equally unsatisfactory. A suitable indicator for the purpose would appear to be Hematoxylin Solution. The presence of neutral Hydrochloride of Quinine is intended to be determined by this test, and this salt is neutral in reaction towards Hematoxylin Solution. If the salt be mixed with an equal weight of Sodium Sulphate, the mixture dissolved in 10 times its weight of hot Water, the liquid neutralised with Ammonia Solution, cooled and set aside at 15.5° C (60° F), the Quinine Sulphate formed is officially required to answer the tests for freedom from other Cinchona alkaloids, given under Quinine Sulphas. The salt should dissolve without change of colour in concentrated Sulphuric Acid, Hydrochloric Acid gas being simultaneously evolved, indicating the absence of readily charred organic impurities. It should yield no red coloration when mixed with a few drops of concentrated Nitric Acid, which the salt is free from Morphine. When ignited with free access of air it should leave no weighable residue.

QUININÆ SULPHAS.

QUININE SULPHATE



FR, SULFATE BASIQUE DE QUININE, GER, CHININSULFAT, IJAR, SOLFATO DI CHININA, SPAN, SULFATO QUINICO BASICO

Light, white, odourless, silky, needle-shaped crystals, with a very persistent bitter taste. It is officially described as the Sulphate of an alkaloid obtained from the Bark of various species of *Cinchona* and *Remyria*, but would have been preferably described, as in the *U.S.P.*, as the Sulphate of the alkaloid Quinine.

The crystals effloresce on exposure to dry air, and yield a salt containing about 2 molecules of Water of crystallisation, these are in turn lost at a temperature of 100° C (212° F), but again reabsorbed on exposure of the dry salt to the air. Such a salt should be made official.

The *B.P.* formula shows 15 molecules of Water of crystallisation, the formula given in the *U.S.P.* shows 7 molecules of Water of crystallisation. According to Cowley there is no doubt that very little of the Quinine Sulphate used for dispensing purposes contains the amount of Water represented by the 15 molecules formula.

The formula of the salt official in the *Fr. Codex* (1908) shows 8H₂O, the basic Quinine Sulphate being stated to crystallise with 8 molecules of Water of crystallisation during the cooling of its hot concentrated aqueous solutions, it contains 72.81 p.c. of Quinine, 11.01 p.c. of Sulphuric Acid and 16.18 p.c. of Water.

The *Fr. Codex* also states that from Absolute Ethylic Alcohol it forms acicular crystals containing 1 molecule of Alcohol of crystallisation which it loses readily.

It should be kept in well closed vessels, preferably in well stoppered glass bottles of a dark amber tint and protected as far as possible from exposure to the light, as in addition to its efflorescent nature in dry air the salt is liable to acquire a yellow or brownish colour when exposed directly to the light

Solubility—About 1 in 800 of Water, 1 in 25 of boiling Water, 1 in 65 of Alcohol (90 p c), 1 in 40 of Glycerin

60 grains require 60 minims of diluted Sulphuric Acid, or 100 minims of diluted Phosphoric Acid for solution in 2 fl oz of Distilled Water

66 grains requires 60 minims of Diluted Nitric Acid for solution in 2 fl oz of Water

Medicinal Properties—In small doses it acts as a most valuable tonic and bitter stomachic. In large doses it has a specific action in malaria, both as a curative and as a prophylactic, in moderate doses it is an antipyretic in influenza and fevers, especially enteric (in which it also acts as an antiseptic), and it is analgesic in supra orbital and other forms of neuralgia. Used as a spray (2 grains to 1 fl oz) in hay fever, contra-indicated during advanced pregnancy and in acute or subacute middle ear disease, in large doses, or if taken frequently, produces temporary deafness. Rectal injections of a strong solution most satisfactory in amoebic dysentery

The best remedy in influenza, also, as a trustworthy prophylactic, 2 grains every morning, the late Sir W. Broadbent, *Pr* '07, i 13, other references to its use as a prophylactic, *B M J E* '95, ii 92, *L* '95, ii 1381

Seems to be really an antitoxin in influenza. If quinine treatment in influenza is persevered with, there will be much less cardiac weakness and fewer serious sequelæ—*Pr* '07, i 153

It retards or arrests the alcoholic, lactic and butyric fermentations, but not the digestive action of Pepsin

In the form of a 1 p c solution in just sufficient Diluted Sulphuric Acid to hold the salt in solution, has been used (*L* '05, i 360) as a powerful curative agent in a large variety of corneal ulcers not amenable to the ordinary routine treatment, the eyes being soaked in the solution for 5 minutes 4 or 5 times a day. As regards the use of Quinine salts in ophthalmic work, it is pointed out (*B M J* '04, i 452) that it has been in employment for twenty years at the Liverpool Eye and Ear Infirmary, and two formulas taken from the Pharmacopœia of this institution are Atropine Sulphate, 4 grains, Quinine Sulphate (neutral), 4 grains, Distilled Water, 1 oz,—and Eserine Sulphate, 1 grain, Quinine Sulphate (neutral), 4 grains, Distilled Water, 1 oz

In whooping cough, *M A* '95, 522, *T G* '94, 126, in cholera nostras—*B M J E* '93, ii 7

As a prophylactic in African fevers—*L* '96, i 219

Combined with Ipecacuanha in dysentery—*P* liv 478, *P J* (8) xlv 1167

10 grains with half its bulk of Tartaric Acid dissolved in 10 minims of Water in the comatose and cerebral forms of remittent fever—*B M J* '99, ii 1474

Quinine and the malarial parasite. As the red blood corpuscle is necessary for the life of the parasite, Quinine, by driving the parasite out of its element, places it under conditions unfavourable and destructive to its development.—*B M J E* '99, ii 68

$\frac{1}{2}$ to 1 dm doses of Ammoniated Tincture in treatment of dengue fever in Canton—*L* '03, i 184

Quinine rash caused by taking not over $\frac{1}{2}$ grain—*T G* '02, 8

Applied as a dressing 1 dm to 8 oz in emulsion with Cod-Liver Oil in tertiary and rheumatic ulcers of the leg—*L* '02, i 443

5 to 10 grains given, dissolved in the acid portion of an effervescent Potassium Citrate mixture, in certain forms of extensive dermatitis—*B M J* '03, i 656, *L* '03, i 785

Six cases of tetanus following the injection of strong solutions of Quinine—*B M J E* '02, i 63

From the results of bacteriological tests Quinine salts seem to be more potent anti-septics than Carbolic Acid or Formaldehyde, and intermediate between these and Corrosive Sublimite—*E M J R* '02, ii 12

In malaria, 2 to 5 grains every 3 or 4 hours, as soon as the diagnosis has been made, yield (*J*) *Exm* (681) better results than large doses at close of or before pyrexia $\frac{1}{2}$ to 1 grain $\frac{1}{16}$ grain Atropine may be given if headache is severe Another '04, ii 1150 is to give 4 doses of 7½ grains, repeated at intervals of ½ hour, in the evening of every third day during the first fortnight of the fever Larger doses—15 to 20 with from 15 to 20 minims of Laudanum have been recommended—*L M J E* '04, ii 1451 In blackwater fever (*B M J E* '04, ii 83) it has been recommended by Koch's method—1 gramme (= 15 grains) on each of 2 consecutive days, at intervals of 10 days

3 to 6 grains every 3 or 4 hours, combined with Ammonium Carbonate in an effervescent mixture form a good prescription in the treatment of puerperal infection Best to begin with Quinine and Calomel, and in the later stages to administer Ferric Perchloride and Magnesium Sulphate—*L* '05, i 1406

In the prophylaxis of malaria a full dose of 10 or 15 grains should be taken on 2 successive days with an interval of 8 or 9 days before the next 2 doses are taken—*L* '05, ii 540

1 drm. of a solution made by dissolving 12 grains of the salt in 30 minims of Distilled Water and 30 minims of dilute Sulphuric Acid, injected (*B M J* '05, ii 724) into each ligament in the treatment of prolapsus uteri

In the pyrexia of pulmonary tuberculosis the only drug which may be tried is Quinine, though it is apt to disturb the stomach It should be given (*J*) *Med Jour* '05, 467) in a single dose of 20 to 30 grains, or 4 or 5 smaller doses, at short intervals

In the leucopenia of cachexial fever and Kala-azar, large doses of Quinine (60 grains daily) combined with red bone marrow have given much better results (*B M J* '05, i 710) than any yet reported by those who deny the value of the drug

2 or 3 grains 3 times daily, combined with the external application of Iodine, have been known to cure an obstinate case of lupus erythematosus in a month Adrenalin may with advantage be combined with the Quinine—*B M J* '05, i 700

In blackwater fever 15 to 18 grains hypodermically at once and 10 to 12 grains 3 times a day for 5 days, and twice a day for 2 following days—*L* '05, ii 599

The value of Quinine in the treatment of blackwater fever has been challenged, and the question whether the fever can be induced by its administration has been discussed A case reported by Dr A D Kitchen in the *South African Medical Record* (*L* '06, ii 820) appears to prove that it can

Dose.—1 to 10 grains = 0.06 to 0.65 gramme

Prescribing Notes.—Given in pills or cachets, also in aqueous solution assisted by the addition of Diluted Sulphuric or Diluted Hydrochloric Acid, 1 minim to each grain, it also dissolves readily in Tincture of Ferric Chloride

One of the most efficacious ways of giving Quinine is in a mixture with Citric Acid, to be taken 3 or 4 times a day It is also given in solution containing Potassium Bicarbonate or in solution with Carbolic Acid It is also given in solution with Hydrochloric Acid to a patient suffering from cinchonism Malt covers the taste well Effervescent Quinine Citrate is also a very palatable form

For disguising the taste of Quinine, when administered to children, Chocolate has been suggested

When a large dose (say 10 grains) is given it is best suspended in Water, the bitterness is not then so intense as when in solution

It is best made into pills with Diluted Glucose

For hypodermic injection see other salts of Quinine, under each of which

the solubilities are given. Of the neutral salts, the Lactate (1 in 1) is the most soluble, of the acid salts, the Acid Hydrochloride (1 in 1).

Quinine is precipitated from aqueous solutions of its salts by alkalis. In the Ammoniated Tincture of Quinine the alkaloid is dissolved by the Alcohol.

Incompatibles All alkalis and their Carbonates, Benzoates, Iodides, and Silicylates, all infusions containing Tannin throw down a Quinine Tannate, which Sulphuric Acid, instead of dissolving, helps to precipitate.

Official Preparations—*Pilula Quininae Sulphatis* and *Tinctura Quininae Ammoniatum*. Used in the preparation of *Pomum* of Quinine Citrus and *Syrupus Ferri Phosphatis cum Quinina* et *Stychnina*.

Not Official—Ammoniated Quinine Capsules, *Elixir Quininae Ammoniatum*, *Mistura Quininae*, *Mistura Quininae cum Ferro*, *Pilula Metallorum*, *Pilula Quininae Sulphatis Composita*, *Atkin's Tonic Pill*, *Pilula Quininae cum Ferro*.

Foreign Pharmacopœias—Official in Austr, Ger, Hung, Jap, Russ, and Swiss (*Chininum Sulfuricum*), Belg (*Sulphas Quininae*), Dan, Norw and Swed (*Sulphas Chinicus*), Dutch (*Sulphas Chinini*), Fr (*Sulfate Basique de Quinine*), Ital (*Solfato di Chinina*), Mex and Port (*Sulfato de Quinina*), Spain (*Sulfato Quinico basico*), U.S. (*Quinina Sulphas*).

Tests—Quinine Sulphate is officially required to lose 11 molecules, equivalent to 11.2 p.c. of Water of crystallisation when exposed to dry air, and a freshly prepared salt should lose, according to the official requirements, 15.2 p.c. of Water when dried at 100° C (212° F). The *USP* states that when exposed to dry air or when heated to 60° C (140° F) it loses 5 molecules, equivalent to 10.3 p.c. of Water of crystallisation, the remaining number of molecules equivalent to 4.1 p.c. being lost at a temperature of 115° C (239° F), indicating a total loss of 14.4 p.c. The *PG* states that the salt shall lose not more than 15 p.c. when heated at a temperature of 100° C (212° F). The *Ph Codex* states that when exposed to the air it rapidly effloresces, losing 6 molecules of Water of crystallisation, equivalent to 12.13 p.c., leaving a salt containing 2 molecules of Water of crystallisation, equivalent to 4.60 p.c. The salt loses the whole of its Water of crystallisation only slowly at 100° C (212° F), but at 115° C (239° F) it becomes rapidly anhydrous. Neither the *BP* nor the *PG* refers to the m.p. of the dried salt. The *USP* states that when dried over Sulphuric Acid it melts at 205° C (401° F). The salt dissolves sparingly in Water, forming a solution which is neutral in reaction towards litmus paper, and which possesses but a slight fluorescence. The *BP* states that the aqueous solution has a bluish fluorescence, the *USP* that the aqueous solution develops a vivid blue fluorescence when acidified with diluted Sulphuric Acid. The *PG* states that the aqueous solution exhibits no fluorescence, but on the addition of a few drops of diluted Sulphuric Acid a blue fluorescence is developed. A solution of the salt affords with Ammonia Solution a white precipitate soluble in excess of the reagent or in Ether. The separated alkaloid answers the tests distinctive of Quinine given under Quinine. When acidified with Hydrochloric Acid the aqueous solution affords with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid.

The more generally occurring impurities are deficiency of Water of crystallisation, readily charred organic impurities, Ammonium

Sulphate and inorganic salts, Morphine, mineral impurities, Cinchonidine, Cinchonine and amorphous Cinchona alkaloids. The *BP*, as above stated, requires that a freshly prepared salt, when dried at 100°C (212°F), should lose 15.2 p.c. of Water. The *USP* requires that the residue remaining on drying 1 gramme of the salt at a temperature of 115°C (239°F) until it ceases to lose weight, should weigh not less than 0.833 gramme, indicating a loss of 16.18 p.c., equivalent to 8 molecules of Water of crystallisation, although the formula given only shows $7\text{H}_2\text{O}$. The *PG* requires that when dried at 100°C (212°F) it shall lose not more than 15 p.c. by weight. The salt should not acquire a faintly yellowish tint when mixed with Sulphuric Acid, indicating the absence of readily charred organic impurities. The *PG* and the *USP* include a test for the absence of Ammonium Sulphate and inorganic salts described in small type below under the heading of Chloroform and Alcohol. No similar test is mentioned in the *BP*. Quinine may be distinguished from Morphine by the Nitric Acid test also described in small type. The *PG* includes a test for Chlorides with Silver Nitrate. 1 gramme of the salt, when ignited with free access of air, should leave no weighable residue. The natural and most probable impurity is Cinchonidine Sulphate, which is generally present to some extent in the commercial Sulphate. The *BP* requires that when tested according to the official method the salt should not yield an appreciable reaction distinctive of Cinchonine, Cupreine, Quinidine or amorphous alkaloid, and that not more than a total of 3 p.c. of impure Cinchonidine crystals should be yielded when the sample is assayed according to the official test, which means about double this amount in the sample. The *BP* includes tests for the alkaloids Quinidine and Cupreine, but they are very unlikely impurities. The test for Cinchonidine and Cinchonine depends upon two principal features, the comparative solubilities of the Sulphates in Water and the relative insolubility of Cinchonidine and Cinchonine in Ether. Quinine Sulphate is soluble 1 in about 800 of Water, Cinchonidine Sulphate 1 in 100 of Water and Cinchonine Sulphate 1 in 70 of Water, and advantage is taken of the comparative insolubility of Quinine Sulphate in cold Water to remove a greater portion of the Quinine by crystallisation. Freese, *loc. cit.* Quinine is readily soluble in Ether, whereas both Cinchonidine and Cinchonine require comparatively large quantities of Ether to effect solution. Opinions differ as to whether it is preferable to effect solution of the Cinchonidine and Cinchonine Sulphates by digesting a weighed quantity of the sample with a limited amount of Water at 60°C (140°F), or to use sufficient Water at a temperature of 100°C (212°F) to dissolve all three Sulphates and to decompose any double Sulphate of the two alkaloids, subsequently cooling to 50°C (122°F). It seems to be generally conceded that simple digestion of the salt with Water at 15.5°C (60°F) does not effect solution of the more soluble salt, owing possibly to the existence of a double Sulphate. When, however, solution of the mixed Sulphates is made in just about sufficient boiling Water to effect solution, the less soluble salt crystallises out almost entirely, leaving the more soluble salt in

solution In the *F*₁ *Coder* method, which is described below, solution is effected at a boiling temperature and the *B P* dissolves a weighed quantity of 4 grammes of the salt in 120 c.c. of boiling Water, and after cooling the solution gradually to 50° C (122° F) with intervals of frequent stirring, filters off the recrystallised Sulphate and evaporates the filtrate to a volume of somewhat less than 10 c.c., which is transferred to a small stoppered flask and shaken when cold with 10 c.c. of Ether and 5 c.c. of Ammonia Solution Any crystals which separate out after the mixture has been allowed to remain at rest in a cool place for not less than 24 hours are collected on a tared filter and dried at a temperature of 100° C (212° F), after previously washing with a little Ether When cool they are weighed and the weight should not amount to more than 0.12 of 1 gramme, indicating not more than 3 p.c. of impure Cinchonidine The *U S P* and the *P C* do not employ the Ether-solubility test, but use that with Ammonia Solution described below The *B P* test has been severely and very adversely criticised It has been shown by Cowley (*P J* '98, 1 412) that Cinchonine and Cupreine are never present in Quinine Sulphate of any known commercial manufacture, moreover, Cupreine occurs in Cupiea Bark (*Remijia pedunculata*), now seldom if ever employed by Quinine manufacturers, and in any case it could only exist in Quinine Sulphate to the extent of a few hundredths p.c. A yield of 3 p.c. of crystals of Cinchonidine (by the *B P* test) really means an admixture of 5.99 p.c. crystallised Cinchonidine Sulphate in Quinine Sulphate answering the *B P* requirements, while the 1885 *B P* stipulated that a Quinine Sulphate should not contain much more than 5 p.c. of Sulphates of other Cinchona alkaloids It would therefore have been better for the Pharmacopœia, failing the insertion of a satisfactory test, to describe that limit of impurity leaving its determination, when necessary, in the hands of those competent to undertake it Paul, who has experimented extensively on the *B P* tests for Cinchonidine, suggests (*C D* '04, 11 429) the following method of procedure—A weighed quantity of 1 gramme of the Quinine Sulphate to be examined is dissolved in 100 c.c. of boiling Distilled Water, the solution after cooling is filtered from the crystallised salt, the filtrate concentrated to 30 c.c. Any further crystals which may have formed are separated by passing the cooled solution through a loose plug of Cotton-Wool fitted in the neck of a funnel, and the volume of the solution is made up to 30 c.c. if necessary, by washing the crystals with a few drops of Water A measured quantity of 5 c.c. of this solution, after adding 5 drops of Ammonia Solution, is shaken with 1 c.c. of Ether in a corked tube, the tube being allowed to remain in a cool place for 1 hour If at the end of that time no crystals are formed in the solution the quantity of Cinchonidine in the 5 c.c. of the solution must be less than 0.004 gramme, and the corresponding quantity of Sulphate in 1 gramme of the salt under examination would not be more than 0.0324 (= 0.004 × 1.35 grammes), or 3.24 p.c. In the more than probable case of crystals being formed in appreciable quantity within a shorter time than 1 hour the amount of the salt

under examination will be more than 3.24 p.c. To ascertain how much more it may be, shake out a volume of less than 5 c.c. with 1 c.c. of Ether, repeat that operation until a difference of 0.5 c.c. of solution, between two experiments, also corresponds to entire absence of crystals in the one instance and the very slight formation of crystals in the other after 12 hours, then take the mean of those two quantities of solution as containing 0.004 gramme of Cinchonidine and calculate the percentage of Sulphate on that basis. Thus for example, if 4 c.c. of solution gave no crystals and 4.5 c.c. only a little after 12 hours, 4.25 c.c. is to be taken as the quantity of Cinchonidine in the calculation, as follows:

$$4.25 \text{ c.c.} \times 0.004 = 30 \text{ c.c.} \times 0.0282 \times 1.35 = 0.038 \text{ gramme in 1 gramme, or } 3.8 \text{ p.c. of Cinchonidine Sulphate in the sample operated upon.}$$

Paul states that the operations requisite in applying the Ether test are extremely simple, and while they admit of being carried out with ease, the results obtainable are not deficient in accuracy. The *USP* and *PG* tests depend upon the amount of Ammonia Solution required to redissolve the precipitate at first formed in a strictly neutral aqueous solution of the salt, from which the greater portion of the Quinine Sulphate has been removed by recrystallisation, so as to produce a clear liquid. The *USP* test is described in small type below under the heading of Ammonia Solution. The *PG* test which differs slightly from the *USP* is virtually as follows. A weighed quantity of 2 grammes of the Quinine Sulphate which has been previously completely dried at a temperature of 40° to 50° C (104° to 122° F) is digested with 20 c.c. of Water for half an hour in a water-bath at a temperature of 60° to 65° C (140° to 149° F) with intervals of frequent shaking, it is then placed in Water at 15° C (59° F.), and allowed to stand for 2 hours with intervals of vigorous shaking. The crystals are separated by filtration through a piece of dry calico of a capacity of about 100 cm. square, the expressed liquid is filtered through a filter prepared from the best filter paper of about 7 cm. diameter. A measured quantity of 5 c.c. of the filtrate having a temperature of 15° C (59° F) is transferred to a dry test-tube, and sufficient Ammonia Solution having a temperature of 15° C (59° F) added to completely dissolve the precipitate at first produced and to produce a clear solution, not more than 4 c.c. of Ammonia Solution should be necessary. The *Ph. Codex* (1908) also employs the Ammonia test for detecting the presence of other Cinchona alkaloids, and for their detection the following method is given. 1 gramme of the official basic Quinine Sulphate is dissolved at a boiling temperature in 3 c.c. of Distilled Water, is allowed to cool to 15° C (59° F) and maintained at that temperature during half an hour, the vessel being immersed in a water-bath maintained at a temperature of 15° C (59° F), and frequently shaken. The liquid is filtered at this temperature, and the 2 following tests are performed on the liquid. (1) A measured quantity of 5 c.c. of the limpid liquid is transferred, by means of a graduated pipette, to an assay tube, and exactly 5 c.c. of a 10 p.c. Ammonia Solution added, care being taken to disturb the flask

as little as possible during the mixing of the liquid. The tube is stoppered and gently inverted several times, the Quinine at first precipitated is redissolved, and a limpid mixture should be obtained, which should remain in this condition during 24 hours. A permanent turbidity or slow deposition of crystals in the previously clear liquid indicates the presence of alkaloids other than Quinine. (2) A measured quantity of 5 c.c. of the original limpid liquid is transferred to a small accurately tared porcelain evaporating basin, evaporated on a water bath at a temperature of 100° C (212° F) until the evaporating basin and its contents no longer show a variation in weight, the residue left on evaporation of the 5 c.c. of liquid should not weigh more than 0.005 of a gramme. The presence of other soluble salts increases the weight of this residue.

The Ammonia test for Quinine Sulphate has been criticised (*C.D.* '05, 1488), and the results of some experiments dealing with the solubility of Quinine in Ammonia are there recorded. With a view of removing several factors tending to invalidate the Ammonia test, the use of a solution of a fixed Hydroxide instead of Ammonia was suggested, Potassium and Sodium Hydroxide were tried, but the Calcium Hydroxide Solution (*B.P.*) was finally chosen, as it was readily made of constant strength, was less liable to impurity, and any decomposition is evident to the eye. In addition to the evidence of solution of the precipitated alkaloid confirmed by the eye, a determination of the Sulphuric Acid radicle is suggested, Phenolphthalein Solution being employed as an indicator of neutrality. It was found that 20 c.c. of a saturated aqueous solution of purified Quinine Sulphate, to which 3 drops of Phenolphthalein Solution were added, required 2 c.c. of Calcium Hydroxide Solution (*B.P.*), whilst 20 c.c. of Cinchonidine Sulphate Solution required 13.7 c.c. of a similar solution, thus a double method of testing the purity of the Sulphate, *eg.*, the solubility of the alkaloids and the quantity of Sulphuric Acid radicle in the aqueous solution is available. The value of the methods was tried with 5 grammes of commercial Quinine Sulphate, each treated with 100 c.c. of Water at 60° C (140° F) for 1 hour, frequently shaking, then cooling to 15° C (59° F), keeping at this temperature for 2 hours, frequently stirring, and then filtering. 20 c.c. of the Quinine Sulphate Solution required 41 c.c. of Calcium Hydroxide Solution (*B.P.*) to form a clear solution, and 2.8 c.c. of a similar solution, when titrated using Phenolphthalein Solution as an indicator. A mixture containing 1 p.c. of Cinchonidine required 45 c.c. of Calcium Hydroxide Solution (*B.P.*) to form a clear solution, and 3.3 c.c. when titrated, using Phenolphthalein Solution as an indicator. A mixture containing 3 p.c. of Cinchonidine required 55 c.c. of the Calcium Hydroxide Solution to form a clear solution, and 3.8 c.c. for titration, whilst a mixture containing 5 p.c. of Cinchonidine required 71 c.c. of the Calcium Hydroxide Solution to form a clear solution, and 4.7 c.c. for titration. In view of the remarks respecting Quindine and Cupreine which appear above, the advantage of retaining tests for the presence of these alkaloids is doubtful, but if their presence is suspected the following brief outlines

of the methods adopted for their detection may be useful. The *BP* test for Quinidine depends upon the precipitation of Quinidine Hydriodide by the addition of Potassium Iodide Solution, precaution being taken to prevent precipitation of amorphous Iodides by the addition of a little Alcohol (90 p c). The major portion of the Quinine Sulphate is removed from a weighed quantity of 1 gramme of the salt by dissolving it in 30 c c of boiling Water, cooling and filtering. A little Alcohol (90 p c) is added to the cold filtrate and sufficient Potassium Iodide Solution to precipitate the Quinidine Hydriodide, which is collected on a tared filter, washed with a little Water, dried till constant in weight, and when cool, weighed, its weight represents about an equal quantity of crystallised Quinidine Sulphate. It is officially required to yield none, or only the slightest traces. Cupreine forms a Sulphate which is sparingly soluble in Water, and this Sulphate will appear with the recrystallised Quinine Sulphate obtained in the first stages of the Cinchonidine test. The *BP* removes it from this crystalline precipitate by adding 6 c c of Ammonia Solution, and shaking with 25 c c of Ether, and this ethereal solution is mixed with the ethereal liquid obtained in the washing of the impure Cinchonidine crystals, and from the mixed ethereal liquids the Cupreine is separated by shaking with 6 c c of Sodium Hydroxide Solution (10 p c), Cupreine forming with an excess of Sodium Hydroxide Solution a definite crystallisable compound which cannot be extracted by Ether. The alkaline layer after separation is shaken with Ether to remove all traces of other alkaloids, and after being exactly neutralised with diluted Sulphuric Acid the Cupreine Sulphate is allowed to crystallise. This involves the presence of a fair quantity of Cupreine, and it would have been preferable to add Ammonium Chloride to the alkaline solution, to extract with Ether, or to neutralise the alkaline liquid with diluted Sulphuric Acid, to add Ammonia Solution and to extract with Ether. In the event of Cupreine being present it will separate out in crystals from the ethereal solution. The *BP* requires the sample to be either free from this alkaloid, or at the most to contain the slightest traces. In testing for Cinchonine and amorphous alkaloids the *BP* dissolves 1 gramme of the Sulphate in 30 c c of boiling Water, precipitating the Quinine and other crystalline alkaloids by the addition of 1 gramme of Sodium Potassium Tartrate, the cold filtered aqueous liquid when evaporated to a small bulk is officially required to yield little or no precipitate with Ammonia Solution. The remarks on the British tests for these latter alkaloids appear above.

Residue.—After ignition the salt should not leave any residue, *BP*, *P.G.* and *USP*.

Chloroform and Alcohol.—1 gramme of the salt, when gently heated to 50° C (122° F) with 7 c c of a mixture of 2 volumes of Chloroform and 1 volume of Absolute Alcohol, should completely dissolve and the solution should remain clear on cooling, *USP* and *P.G.*, the latter stating a temperature of from 40° to 50° C (104° to 122° F).

Sulphuric Acid.—Sulphuric Acid should impart to the salt not more than

a faintly yellowish tint, *U S P*, the salt should scarcely be coloured when moistened with Sulphuric Acid, *P G*

Nitric Acid—Nitric Acid should not produce a red colour, *U S P*, the salt should scarcely be coloured when moistened with Nitric Acid, *P G*

Silver Nitrate—An aqueous solution of the salt, after acidifying with Nitric Acid, should not be affected by *T S* of Silver Nitrate, *P G*

Ammonia Solution—The Quinine Sulphate is dried in a porcelain dish on a water bath for 2 hours at 50° C (122° F)

Transfer a weighed quantity of 1.8 grammes of the dried residue (which should be neutral or slightly alkaline to Litmus paper) to a dry test tube and agitate it with 20 c c of Distilled Water for half an hour at 65° C (149° F) and then allow it to cool to 15° C (59° F), and keep the temperature at 15° C (59° F) for 2 hours, shaking the test tube occasionally. Filter the liquid, transfer 5 c c of the filtrate to a test tube, and carefully add 7 c c of Ammonia Water (which must be of official strength and have the temperature of exactly 15° C (59° F) and must be all added at once). A clear liquid should result. If the temperature during the maceration has been 16° C (60.8° F) 7.5 c c of Ammonia Water may be added. If 17° C (62.6° F) 8 c c may be added (limit of allowable foreign Cinchona alkaloids), *U S P*

Preparations

PILULA QUININÆ SULPHATIS PILL OF QUININE SULPHATE

Triturate 30 grains of Quinine Sulphate with 1 grain of Tartaric Acid, and add them to the previously mixed Glycerin, 4 grains, and Tragacanth, 1 grain

Dose—2 to 8 grains = 0.13 to 0.52 gramme

TINCTURA QUININÆ AMMONIATA AMMONIATED TINCTURE OF QUININE

Quinine Sulphate, 175 grains, Solution of Ammonia, 2 fl oz, Alcohol (60 p c), 18 fl oz (about 1 grain in 55 minims)

When first made the Tincture usually deposits a little, so it is better to allow a day or two to elapse before filtering

Dose—½ to 1 fl drm = 1.8 to 3.6 c c

When mixed with Water the Quinine is precipitated in a fine state of division, but the particles soon aggregate and adhere to the sides of the glass, therefore this preparation should not be prescribed in mixtures unless Mucilage of Acacia be used to suspend the Quinine

When prepared with Ammonium Carbonate instead of Liquor the Tincture does not precipitate so badly, and it may be diluted with Water saturated with Carbonic Acid without any precipitation at all

Tests—Ammoniated Tincture of Quinine possesses a sp gr of 0.9225 to 0.9230, it contains about 1.8 p c w/v of total solids and about 54 p c w/v of Absolute Alcohol. 10 c c of the Tincture requires for neutralisation about 5.5 c c of Normal Volumetric Sulphuric Acid Solution, using Cochineal Solution as an indicator of neutrality

Not Official

AMMONIATED QUININE CAPSULES—Quinine Sulphate, 60 grains, Ammonium Carbonate (powdered finely), 100 grains, Soft Paraffin and Liquid Paraffin, *q s* to make a thin paste and fill 100 capsules. Each capsule represents about 30 minims of Ammoniated Tincture of Quinine

ELIXIR QUININÆ AMMONIATUM —Quinine Sulphate, 1, Ammonium Carbonate, 3, Alcohol, 25, Solution of Carmine, 0.25, Elixir of Orange, 50, Distilled Water, *q s* to produce 100 — *B P C*

MISTURA QUININÆ —Quinine Sulphate, 1 grain, Diluted Sulphuric Acid, *q s*, Distilled Water, to 1 fl oz — *London Ophthalmic*
Quinine Sulphate, 1 grain, diluted Sulphuric Acid, 1 minim, Tincture of Orange, 10 minims, Water, to 1 fl oz — *St Thomas's*

This has been incorporated in the *B P C*

MISTURA QUININÆ CUM FERRO —Quinine Sulphate, 1 grain, Solution of Ferric Chloride, 10 minims, Water, to 1 fl oz — *St Thomas's*

This has been incorporated in the *B P C*

PILULÆ METALLORUM —Quinine Sulphate, 1 grain, Reduced Iron, 1 grain, Strychnine (alkaloid), $\frac{2}{10}$ grain, Arsenic Trioxide, $\frac{1}{10}$ grain, in one pill — *U S N F*

Note A similar combination is known under the name of Aitken's Tonic Pills —

Quinine Sulphate, 1 grain, Reduced Iron, $\frac{3}{4}$ grain, Strychnine (alkaloid), $\frac{1}{10}$ grain, Arsenic Trioxide, $\frac{1}{10}$ grain, in one pill — *U S N F*

This has been incorporated in the *B P C* as follows —

Pilulæ Quininae Sulphatis Compositæ *Syn* Aitken's Tonic Pills — Quinine Sulphate $\frac{1}{2}$ grain, Reduced Iron, $\frac{3}{4}$ grain, Arsenious Anhydride, $\frac{1}{10}$ grain, Strychnine, $\frac{1}{10}$ grain, Extract of Gentian, *q s*, in 1 pill — *B P C*

Aitken's Tonic Pill —Quinine Sulphate, 1 grain, Reduced Iron, $\frac{3}{4}$ grain, Arsenious Anhydride, $\frac{1}{10}$ grain, Strychnine, $\frac{1}{10}$ grain, Extract of Gentian, *q s*
Pharm Form

PILULA QUININÆ CUM FERRO —Quinine Sulphate, 1 grain, Ferrous Sulphate, 1 grain, Extract of Gentian 3 grains, in each — *St Thomas's*

Quinine Sulphate, 1 grain, Exsiccated Ferrous — — — — — each pill — *St Thomas's*

This has been incorporated in the *B P C*

RESINA.

RESIN

FR, COLOPHANE, GER, KOTOPHONIUM, ITAL, COLOFONIA, SPAN, COLOFONIA

A translucent, pale amber-coloured, brittle solid, having a turpentine odour. Readily reduced to powder. It is officially described as the residue from the crude Oleo-Resin of various species of *Pinus*, after the Oil of Turpentine has been removed by distillation.

Solubility —In almost all proportions of Alcohol (90 p.c) Ether, and Oil of Turpentine, also in hot Olive Oil.

Medicinal Properties —Antiseptic, and slightly stimulant. It is an ingredient of plasters used for superficial wounds. The ointment forms a stimulating dressing for indolent ulcers and sores. Never used internally.

Official Preparations —Emplastrum Resinae and Unguentum Resinae. Used in the preparation of Emplastrum Calefaciens, Emplastrum Cantharidis, Emplastrum Menthol, Emplastrum Picis, Emplastrum Plumbi Iodidi, Emplastrum Saponis.

Not Official —Resina Carbolicæ, Resina Carbolicata.

Resin Plaster is contained in Emplastrum Belladonnae, Emplastrum Opii, also in Emplastrum Calefaciens.

Foreign Pharmacopœias—Official in all as Colophonium, Span (Colofonia and Resina commune) and U S (Resina)

Descriptive Notes—The Resin of commerce is met with in various grades, from the nearly black Colophony to the water-white or almost colourless, transparent kind. The official variety apparently agrees with the characters of the grade known in trade as Amber Resin. It is transparent, amorphous and very brittle, the freshly fractured surface is shiny and slightly concave, with a faintly terebinthinate odour. The Resin of commerce varies in the amount of Turpentine Oil that it retains. The 'water-white' and 'window-glass' Resins are useful for colourless varnishes. The yellow opaque Resin is made by stirring Water into the Resin after distillation of the Oil of Turpentine, but it loses Water and becomes translucent when heated. Powdered Resin should not cohere into masses.

Tests—Resin has a sp gr of 1.07 to 1.085, the *USP* states 1.070 to 1.080, the *PG* does not refer to a sp gr. When heated it melts, when strongly heated it evolves heavy white vapours possessing an aromatic odour, and when ignited burns readily with a yellow flame, emitting a dense sooty smoke. It dissolves readily and completely in Alcohol (90 p c), Benzol, Carbon Bisulphide and Ether. The *USP* states that it is soluble in Acetic Acid (36 p c w/w), Alcohol (94.9 p c), Benzene, Carbon Bisulphide, Ether, fixed or volatile Oils, and in Potassium or Sodium Hydroxide Solutions. The *PG* states that it dissolves slowly in 1 part of Alcohol (90 p c), and in 1 part of Acetic Acid (96 p c w/w), also in Sodium Hydroxide Solution (15 p c w/w). The Acid value varies from 150 to 185, the Ester value from 0 to 12, the Saponification value from 179 to 193. The *USP* states the Acid value should not be less than 150, the *PG* 151.6 to 179.7. Neither the *USP* nor the *PG* includes an Ester or Saponification value. Neither an Acid, Ester nor Saponification value is included in the *BP*. The presence of Turpentine Oil may be detected by the solubility of the Resin in Alcohol (90 p c). When ignited with free access of air it should burn leaving no weighable residue, indicating the absence of mineral impurities.

Acid Value—A weighed quantity of 1 gramme of the Resin is dissolved in a sufficiency of Alcohol (94.9 p c), a few drops of Phenolphthalein Solution added, and the mixture titrated with Normal Volumetric Potassium Hydroxide Solution. The number of c c of Normal Volumetric Potassium Hydroxide Solution consumed multiplied by 0.05371 indicates the number of grammes of Potassium Hydroxide, and this figure expressed in mg indicates the Saponification value of the Resin, which in this case should be not less than 150, *USP*.

A weighed quantity of 1 gramme of the Resin is dissolved in 25 c c of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution, and after the addition of 10 drops of Phenolphthalein Solution the excess is titrated with Semi-normal Volumetric Hydrochloric Acid Solution, from 18.6 to 19.6 c c should be necessary to neutralise this excess, *PG*.

Preparation

EMPLASTRUM RESINÆ RESIN PLASTER *BP Syn*—ADHESIVE PLASTER

Resin, 4, Lead Plaster, 32, Hard Soap, 2 (1 in 9½)

Now made with Hard Soap instead of Curd Soap

Foreign Pharmacopœias—Official in Aust, Lead Plaster 10, Wool Fat 1, Yellow Wax 1, Turpentine 1, Colophonium 1, Dammar 1, Belg and Swiss, Lead Plaster 80, Elemi 5, Yellow Wax 5, Colophonium 5, Turpentine 5; Dan and Swed, Lead Plaster 8, Colophonium 2, Dutch, Lead Plaster 70, Gum Elastic 10, Wool Fat 20, Gor, Lead Plaster 40, Solid Paraffin 2 5, Liquid Paraffin 2 5, Colophonium 35, Dammar 10, Caoutchouc 10 Petroleum Benzine 75, Hung, Lead Plaster 400, Purified Colophonium 100, Turpentine 25, Ital, Lead Plaster 40, Burgundy Pitch 7, Yellow Wax 3, Mex, Lead Plaster 100, Yellow Wax 10, Dammar 10, Colophonium 10, Turpentine 10, Norw, Lead Plaster 8, Yellow Wax 1, Mastic 1, Russ, Litharge 11, Olive Oil commune 10, Lard 10, Colophonium 8 5, and U S, Lead Plaster 96, Rubber 2, Petrolatum 2, all (Emplastrum Adhæsivum) Jap (Emplastrum Resina), Lead Plaster 80, Yellow Wax 6, Resin 14, Span, Emplastro de Resinas Aglutinante, Lead Plaster 60, Olive Oil 75, Turpentine 75, Yellow Wax 90, Elemi 180, Pine Resin 570

UNGUENTUM RESINÆ. RESIN OINTMENT *NO Syn -*
BASILICON OINTMENT

Resin, in powder, 8, Yellow Beeswax, 8, Olive Oil (by weight), 8, Lard, 6 (1 in 3½)

Olive Oil and Lard used in place of Almond Oil and Simple Ointment, and the quantity of Beeswax increased

Foreign Pharmacopœias—Official in Aust (Unguentum Basilicum), Yellow Wax 16, Olive Oil 36, Colophonium 12, Suet 12, Turpentine 12, Pitch 12, Dutch (Unguentum Resinosum Flavum), Yellow Wax 18, Colophonium 8, Sesame Oil 70, Turpentine 4, Fr (Pommade de Styriax), Purified Liquid Storax 16, Colophonium 29, Purified Elemi 16, Yellow Wax 16, Olive Oil 23, Ger, (Unguentum Basilicum), Olive Oil 9, Yellow Wax 3, Colophonium 3, Suet 3, Turpentine 2, Mex (Unguento Amarilla), Yellow Wax 6, Colophonium 5, Suet 4, Aceite 12, Norw (Unguentum Basilicum Nigrum), Colophonium 12, Yellow Wax 12, Pitch 12, Suet 12, Turpentine 12, Olive Oil 40, Port (Unguento de Resina), Yellow Wax 25, Resin 25, Oleo de Amendoim 50, Span (Unguento de Altea), Turpentine 50, Althæa Root 100, Water 100, Yellow Wax 160, Pine Resin 160, Olive Oil 750, Swed (Unguentum Terebinthinæ Resinosum), Colophonium 15, Suet 15, Turpentine 10, Yellow Wax 15, Olive Oil 45, Swiss (Unguentum Resinosum), Colophonium 9, Turpentine 9, Yellow Wax 17, Olive Oil 65, U S (Ceratum Resinæ), Resin 35, Yellow Wax 15, Lard 50, also (Ceratum Resinæ Compositum), Resin 225, Yellow Wax 225, Prepared Suet 300, Turpentine 115, Linseed Oil 135

Not Official

RESINA CARBOLICA (R D H)—Resin, 4 parts, Carbolic Acid 4 parts; Chloroform, 3 parts Dissolve and filter

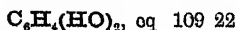
Resina Carbolisata—Carbolic Acid, 3 5, Resin, in powder, 4 5, Chloroform, 2 0—*B P C*

Not Official.

RESORCINUM.

RESORCIN

METADIOXYBENZOLIN RESORCINOL



Fr, Résorcine, GER, RESORCIN, ITAL, RESORCINA, SPAN, RLSORCINA

White, or nearly white, glistening, needle-shaped, or prismatic crystals, having a peculiar characteristic odour, and sweetish, pungent taste. It may be obtained by the destructive distillation of Brazilin, or by fusing Sodium Benzol-sulphonate with Sodium Hydroxide

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from exposure to the light, as it has a tendency to acquire a pinkish tint. It is described in the *U S P* as a diatomic Phenol and under the title of Resorcinol. On this account care should be taken not to confound it with the proprietary preparation known also under the name of Resorcinol, which is described below and which is a mixture of equal parts of Iodoform and Resorcin.

Solubility—4 in 3 of Water, 4 in 3 of Alcohol (90 p c), 1 in 1 of Glycerin, 1 in 1 of Ether, 1 in 22 of Olive Oil.

Medicinal Properties—A powerful antiseptic. It is also antipruritic, but it is very depressing to the heart and is dangerous. As a *spray* (1 or 2 p c) in diphtheria and whooping cough, *P* liv '81, 5 to 10 p c solutions in Glycerin, 5 to 10 p c ointments in skin diseases, *B M J* '83, 1 435, *L* '83, 1 570, '90, 1 1317, '91, 1 505, 1165, *T G* '90, 279. In the rosacea, *P* li 380, in pruritus, *M J* '95, 136, in diarrhea and gastric affections, and as a local germicide and stimulant in ulcers and in pharyngitis and chronic rhinitis, *J B T* '94, 463, in leucoplakia, *T G* '95, 141. Untoward effects when administered internally as a powder, *L* '95, 1 779 436, internal administration of 3 grams taken every 4 hours, followed by appearance of Phenol sulphates in the urine and kidney disturbance—*B M J* '01, 1 1266. A watery solution of about 5 grains to the oz, combined with a little alkali employed as a spray, is recommended (*B M J* '05, 1 1680) in the treatment of common cold in the head, and as a mouth and nasal spray in influenza—*L* '07, 1 152.

Dose—1 to 5 grains = 0.06 to 0.32 gramme.

Prescribing Notes—It is frequently prescribed in hair lotions and washes for removing dandruff, but when mixed with an alkali, e.g., Potassium Carbonate, the solutions rapidly darken in colour and acquire a strong green fluorescence, and such lotions frequently produce an unpleasant colouring effect on the hair which, once produced, is somewhat difficult to remove. It is also incompatible with Spiritus Aetheris Nitrosi.

Antidotes—White of Egg, wash out the stomach with Soda or Saccharated Lime, well diluted, stimulants, Atropine, Amyl Nitrite—*Murrell*.

In large doses it produces profuse perspiration, flushing of the face, and giddiness. Dr. Murrell describes a case of poisoning by 2 drms of it which nearly proved fatal—*M T* '81, 1 487.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Russ., Span., Swed., Swiss and U.S.

Tests—Resorcin melts, when pure, at 118° to 119° C (244.4° to 246.2° F), volatilising completely at a higher temperature. The *U S P* gives 109° to 111° C (228.2° to 231.8° F), the *P* Codex (1908) 119° C (246.2° F). It boils at 276° C (526.8° F). The *U S P* gives the boiling point 276.5° C (529.7° F). It dissolves readily in Water, forming a clear solution, which is neutral in reaction towards Litmus paper, and which yields, on the addition of Ferric Chloride T.S., a deep violet colour, which changes to a brownish yellow on the addition of Ammonia Solution. On gently warming 0.05 of a gramme with 0.1 of a gramme of Tartaric Acid and 10 drops of Sulphuric Acid a deep (arming red) liquid is obtained. 0.5 of a gramme, when mixed with 2 cc of Formalin Solution and 2 cc of Potassium Hydroxide Solution (5 p c), and the mixture heated to boiling, a deep red coloration is gradually developed.

The more generally occurring impurities are Catechol, Quinol, empyreumatic bodies, Phenol, and mineral impurities. Catechol gives, with Ferric Chloride T.S., a dark green coloration changing to violet red on the addition of Ammonia Solution, which distinguishes it from Resorcin. Quinol forms green crystals of Quinhydrone on the addition of Ferric Chloride T.S., changing to yellow, sparingly soluble Quinone on the addition of an excess of the reagent. It also yields a white precipitate with neutral Lead Acetate Solution, whereas a solution of Resorcin yields no precipitate. Lead Subacetate Solution produces, however, a white precipitate in an aqueous solution of the salt. The concentrated aqueous solution should be colourless, indicating the absence of empyreumatic bodies,

—Gruy's

This has been incorporated in the *B P C*

This has been incorporated in the *B P C*

This has been incorporated in the *B P C*

This has been incorporated in the *BPC* under the title *Unguentum Resorcini cum Amylo*, with the synonym as above

RESORCIN PLASTER MULL (*Unna*)—Contains $\frac{3}{4}$ grain to the square inch

Spiritus Resorcini	Capillaris, Spiritus Capillorum, Lotion
Resorcin	Castor Oil, 2 50, Cologne Spirit, 20,
Alcohol	

This has been incorporated in the *BPC* as follows — Resorcin, 12 50, Glycerin, 12 50, Hydrous Wool Fat, 25, Soft Paraffin, *q s* to produce 100

UNGUENTUM RESORCINI COMPOSITUM—Resorcin, 8, Distilled Water, 8, Oil of White Buch, 8, Oxide of Zinc, 8, Vaseline, 32, Anhydrous Lanolin, 32. Dissolve the Resorcin in the Water and mix with the other ingredients.—*Bournemouth Formulary*

This has been incorporated in the *B P C* with slight modification.

TRIBROMO-RESORCIN—Minute, white or whitish crystals. It is a powerful antiseptic and bactericide.—*PJ* '99, 11 216

Resorcin Monacetat (Euresol) — A transparent yellow viscous mass, readily soluble in Acetone

RESORCIN CAMPHOR—A liquid obtained by heating together equal parts of Camphor and Resorcin. Is superior to mercurial ointment in removing pediculi.—*P J '96*, 1 229, 826

RESORCINOL—Obtained by melting together equal volumes of Resorcin and Iodoform. It is a red-brown powder, partially soluble in Water, soluble in Ether. Has been introduced as a substitute for Iodoform as a dressing—*PJ '96*, 1 446

ANUSOL (Bismuth Iodo-resorcin-sulphonate) is employed in suppository form in the treatment of piles.—*P J '96*, ii 378

FLUORESCHEIN (Resorcinol-Phthalein Anhydride) — An amorphous yellow-red powder, almost insoluble in Water, in Alcohol (90 p c), and in Ether.

Prepared by the action of Phthalic Anhydride on Resorcin. It dissolves readily in solutions of the alkali hydroxides, *e.g.*, Sodium Hydroxide, forming **Sodium Fluorescein**, a yellowish or greyish red powder readily soluble in Water. In the form of a 2 per cent solution it has been used for staining the denuded spots of the cornea, and has thus been found useful in the diagnosis of corneal ulcers.

LIQUOR FLUORESC EIN Fluorescein 9.5 grms, Sodium Bicarbonate 12 grms, Distilled Water 1 fl oz *London Ophthalmic*

Not Official

RHAMNI FRANGULÆ CORTEX

Syn—CORTEX FRANGULÆ

The dried Bark of *Rhamnus frangula* L. Collected from the young Trunk and from the lower Branches and kept at least one year before being used.

Official in *L P* 1885, but not in *L P* 1899

Medicinal Properties Similar to those of *Rhamnus Purshiana*. A laxative or purgative for delicate constitutions and the aged.

A solid **Extract**, dose, 15 to 60 grms, was official in *L P* '85, and is now in Dutch, Russ and Swed. A **Fluid Extract**, dose, 1 to 4 fl drms, was official in *L P* '85, and is now in Dan, Dutch, Fr Ger, Norw, Russ, Swed, Swiss and U S. Swed. also has a **Syrup**. *Fructus Rhamni Cathartica* is official in Belg. and Ger., also the **Syrup**.

Descriptive Notes The bark is found in commerce as a waste product, being derived from the wood known in trade as 'dogwood', which is imported from Holland for use as gunpowder. The thin bark of the younger trunks and branches is preferable for use in medicine, the thicker bark of old trees being very bitter and nauseous. It requires to be kept a year before being used like that of *R. Purshiana*, the recently collected bark being liable to produce colic, nausea, and vomiting. The dried bark is in the form of thin quills of a dark greyish-brown or greenish black colour externally, and has a brownish yellow inner surface. It should not exceed $\frac{3}{8}$ inch (1 mm.) in thickness. The outer surface is covered with numerous elongated, transverse, whitish marks (lenticels), when the epidermis is rubbed with the nail, a purplish red or dull crimson layer is seen beneath, which forms a characteristic feature of the bark. It does not contain stone cells.

Preparations

EXTRACTUM RHAMNI FRANGULÆ—*Rhamnus Frangula* Bark, in No. 40 powder is percolated with Proof Spirit (Alcohol 57 per cent) until exhausted, the liquor is evaporated by a water bath to an extract—*L P* 1885.

The *L P C* employs the Bark in No. 20 powder, and exhausts by percolation with Water.

EXTRACTUM RHAMNI FRANGULÆ LIQUIDUM—Boil 16 of the Bark in successive quantities of Water, evaporate the liquors to 12, and when cold add 1 of Rectified Spirit, filter, and add Water *q s* to make 16—*L P* 1885.

FLUIDEXTRACTUM FRANGULÆ—Percolate 100 of Bark with a mixture of 50 of Alcohol (95 per cent), with 80 of Water, reserve the first 80, and evaporate the remainder to a soft extract, which dissolve in the reserved portion and make up to 100. *U S P*.

Average Dose—15 minims = 0.9 cc.

This has been incorporated in the *B P C*.

RHEI RADIX.

RHUBARB ROOT

FR, RHUBARBE DE CHINE, GER, RHABARBER, ITAL, RABARBARO;
SPAN, RUIBARBO

Though called Rhubarb Root, it really consists chiefly of the erect Rhizomes of *Rheum palmatum*, L, *R. officinale*, Baill., and probably other species, collected in North-Western China and Tibet

Medicinal Properties.—Cathartic and astringent, the purgative effect precedes the astringent, and therefore Rhubarb is useful in diarrhoea when an aperient is indicated. Stomachic tonic in small doses. Given in dyspepsia, and in occasional but not in chronic constipation. It is non-irritant, suitable for delicate constitutions, and for increasing the effect of other cholagogues and cathartics, useful in hæmorrhoids. It is frequently combined with an antacid or carminative.

Dose.—3 to 10 grains = 0.2 to 0.65 gramme, for repeated administration, for a single administration, 15 to 30 grains = 1 to 2 grammes

Prescribing Notes.—May be given in cachets, pills, mixtures, or Compressed Tablets. The compound powder is also prescribed in cachets, capsules, etc.

4 grains of Powdered Rhubarb and 1 minim of 'Dispensing Syrup' make a nice pill. Sodium Bicarbonate in equal weight with Powdered Rhubarb counteracts the astringency, and covers the taste, the addition of Peppermint Water still further hides it, or 1 drop of Oil of Peppermint, 30 grains of Sugar, will disguise the taste of 15 grains of Powdered Rhubarb, or 1 drop of Oil of Caraway, 30 grains of Sugar, and 10 grains of Powdered Rhubarb, make a good draught with Water to 1½ fl oz.

Official Preparations.—Extractum Rhei, Infusum Rhei, Liquor Rhei Concentratus, Pilula Rhei Composita, Pulvis Rhei Compositus, Syrupus Rhei, Tinctura Rhei Composita.

Not Official.—Elixir Rhei, Extractum Rhei Compositum, Fluidextractum Rhei, Infusum Rhei Concentratum, Pulvis Rhei cum Magnesia, Mistura Rhei cum Soda, Pilula various, Pulveres various, Tinctura Rhei Aquosa, Vinum Rhei, Purgatum and Rumein.

Foreign Pharmacopœias.—Official in all

Descriptive Notes.—The official Rhubarb Root is attributed to *Rheum palmatum*, Linn., *Rheum officinale*, Baill., and probably other species (*Rheum palmatum* var. *Tanguticum* Max, U.S.P.), and is stated to be collected in China and Tibet. The official description covers several varieties. The Chinese Rhubarb Root of commerce occurs either in transverse sections or split longitudinally, it varies in length and diameter, but averages 3 to 4 in. long and 2 to 3 in. broad, although pieces are sometimes met with as much as 6 or 8 in. long and broad in proportion. The outer surface is convex from having been scraped, or sometimes angular from having been sliced, and presents here and there stellate markings, due to the transversely cut medullary rays of lateral buds or of roots. The outer surface is brownish-yellow, but the broken surface pinkish-brown or greyish-brown, the substance is tough and hard, and gritty.

when chewed. The taste is bitter and astringent, and the flavour disagreeable. Although the root of *Rheum officinale* is cultivated in England, only that collected in China and Tibet is official. The Shensi Rhubarb is considered the best, that of Szechuen and Kansuh are less valuable. The drug, which is produced chiefly in the provinces of Shensi, Szechuen and Kansuh, finds its way to Europe usually via Hankow, Shanghai and Canton, although the drug is also produced in other provinces and in Manchuria. The Shensi Rhubarb exhibits, mostly on the lighter and less compact pieces, a rhomboidal network of whitish veins, but none of the known species of Rheum possess this character, so that it is evidently derived from an undescribed species. In commerce the pieces formed by dividing the rootstock longitudinally are known as 'flats' and those cut transversely as 'rounds'. 'High dried' Canton Rhubarb has usually been dried by artificial heat, and when prepared in this way the pieces are apt to become rotten in the centre, hence, as a test of quality, transversely broken pieces are usually exhibited at the drug sales. The hardest and heaviest pieces are usually selected for trimming, which is done by filing. The English cultivated Rhubarb, prepared from *R. rhaponticum*, L., is less gummy than the Chinese and is less active as a purgative, but it gives a brighter yellow powder. The *R. officinale* grown in England is distinguishable from that of *R. rhaponticum* by its larger size and by the dark or blackish red veins traversing it as compared with the reddish-brown veins of the latter species, which usually form more or less parallel lines on the longitudinal section, and radiate lines on the transverse one. In the Chinese Rhubarb, except in the very inferior kinds, the bark is entirely removed, the holes, through which the string is used for suspending the roots in drying, are dark coloured and irregular, and the outer surface of the pieces is convex. In English Rhubarb, which is always dried by stove heat, the outer surface is denuded of the outer layer only, and is always more or less shrunken and irregular, the internal portion is soft and can be easily indented, and the holes, when present, are round and have fresh edges, having been made with a rat-tail file to imitate the Chinese drug. The larger pieces of the English Rhubarb are mostly exported to the United States, the lateral roots, known as 'stick Rhubarb', are sold at a cheap rate by herbalists. The powdered English Rhubarb, apparently from containing or absorbing more moisture, is liable to turn pink when mixed with Magnesia to form Gregory's Powder. In making aqueous preparations of Rhubarb, the use of pieces cut small, rather than coarsely powdered, gives brighter preparations, which are more easily filtered. The raphides being more abundant in Chinese Rhubarb than in English, the percentage of ash affords some indication of the kind used for the powder, that of Chinese Rhubarb yielding according to Hanbury 12.9 to 13.87 p.c. of ash, one sample, however, yielding as much as 43.27 p.c. English Rhubarb afforded 10.90 p.c. of ash.

The *P.G.* mentions that the sphaeraphides measure up to 0.1 mm and the roundish starch grains from 0.003 to 0.018 mm. (0.005 to

0.020 mm USP) which are either simple or grouped two or three together and have an evident hilum

Tests—The ash of Rhubarb Root varies from 7 to 12 p c, 12 samples of picked root examined in the author's laboratory yielded from 4.1 to 21.5 p c, with an average of 9.85, 14 samples of the powder yielded from 6.7 to 12.1 p c, with an average of 8.97.

Preparations

EXTRACTUM RHEI. EXTRACT OF RHUBARB

Rhubarb Root, exhausted with Alcohol (60 p c), and the resulting liquor evaporated to dryness

Dose.—2 to 8 grains = 0.13 to 0.52 gramme

Official in Aust, Dutch, Ger, Jap, Norw, Swed, Swiss and U S, with Spirit and Water mixed, Belg, with Alcohol (60 p c), Dan, with Alcohol (70 p c), Fr, Hung, Ital, Mex, Port, Russ and Span, with Water

Mex and U S have also a **Fluid Extract**, 1 in 1, Belg, Fluid Extract contains g 30 p c of dry residue

INFUSUM RHEI. INFUSION OF RHUBARB

Rhubarb Root, in thin slices, 1, boiling Distilled Water, 20
Infuse 15 minutes, strain (1 in 20)

Now 1 in 20 instead of 1 in 40, and the time is reduced

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 c c

Foreign Pharmacopœias—Official in Belg, Fluid Extract 10, Potassium Carbonate 1, Cinnamon Water 89, Dan, Rhubarb 125, Sodium Carbonate 25, Distilled Water 2000, Concentrated Spirit 125, Cinnamon Water 150, Ital, Rhubarb 3, Sodium Carbonate 1, Water 50, Norw, Rhubarb 25, Sodium Bicarbonate 3, Distilled Water 170, Cinnamon Water 30, Swed, Rhubarb 10, Sodium Carbonate 2, Alcohol (64 p c), g s, Distilled Water, g s to make 100, all Infusum Rhei Alcalinum Fr (Tisane de Rhubarbe), Rhubarb 5, Distilled Water 1000, Span (Infusion de Rhubarbe), Rhubarb 2, Water 50
See also Tinctura Rhei Aquosa, p 1017

LIQUOR RHEI CONCENTRATUS. CONCENTRATED SOLUTION OF RHUBARB

10 of Rhubarb Root, percolated with Alcohol (20 p c), to yield 20
(1 in 2)

Dose.— $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 c c

Tests.—Concentrated Solution of Rhubarb has a sp gr of 1.020 to 1.030, it contains about 12 p c w/v of total solids and about 18 p c w/v of Absolute Alcohol

PILULA RHEI COMPOSITA. COMPOUND RHUBARB PILL

Rhubarb Root, 3 oz, Socotrine Aloes, 2 $\frac{1}{2}$ oz, Myrrh, 1 $\frac{1}{2}$ oz, Hard Soap, 1 $\frac{1}{2}$ oz, Oil of Peppermint, 1 $\frac{1}{2}$ fl drm, Syrup of Glucose (by weight), about 2 $\frac{3}{4}$ oz.

Dose.—4 to 8 grains = 0.26 to 0.52 gramme

5 grains = about $\frac{1}{2}$ grain of Rhubarb and 1 grain of Aloes

Official in Jap, Swiss and U S

PULVIS RHEI COMPOSITUS. COMPOUND POWDER OF RHUBARB *B P Syn*—GREGORY'S POWDER

Rhubarb Root, 2, Light Magnesia, 6, Ginger, 1 (1 in 4 $\frac{1}{2}$)

If a less bulky powder be desired, Heavy Magnesia is officially permitted to be employed

Dose —20 to 60 grains = 1.3 to 4 grammes

Foreign Pharmacopœias— Official in Austri, Magnesium Carbonate 4, Rhubarb 2, Elæosaccharum Fennicul 1, Dan, Norw and Swed, Magnesium Carbonate 1, Rhubarb 1, Elæosaccharum Fennicul 1, Ger and Jap, Magnesium cum Carbonate 10, Rhubarb 3, Elæosaccharum Fennicul 7, Russ, Magnesium Carbonate 1, Rhubarb 1, Elæosaccharum Fennicul 2, all Pulvis Magnesie cum Rheo. Jap has also Pulvis Rhei Compositus, Rhubarb 2, Burnt Magnesia 6, Ginger 1, Syris (Pulvis Magnesie Compositus), Magnesium Carbonate 5, Rhubarb 2, Elæosaccharum Fennicul 3, U.S. (Pulvis Rhei Compositus), Rhubarb 25, Magnesium Oxide 65, Ginger 10.

Pulvis Rhei cum Magnesia *Syn* Improved Gregory's Powder — Rhubarb Root, in powder, 22, Magnesium Carbonate, 60, Ginger, in powder, 11
P.P.C.

SYRUPUS RHEI. SYRUP OF RHUBARB

Rhubarb Root, 1, Coriander Fruit, 1, Refined Sugar, 12, Alcohol (90 p.c.), 4, Distilled Water, 12, should yield about 20 by weight

L.P. directions are not very satisfactory. It is more convenient to make a (1 in 4) fluid Extract of Rhubarb with Alcohol (60 p.c.), evaporate 8 fl oz of the fluid Extract to 3 fl oz, mix this and 5 minims of Oil of Coriander with 24 oz of Sugar, and add Water to make the weight 40 oz, dissolve in the cold, and filter.

Dose $\frac{1}{2}$ to 2 fl drim = 1.8 to 7.1 cc

Foreign Pharmacopœias— Official in Austri, Rhubarb 10, Borax 2, Spirits of Wine diluted 10, Water 90, after 24 hours filter and to 10 of filtrate add 16 of Sugar. Dutch, Rhubarb 30, Sodium Carbonate 3, Water 150, to 150 of liquid add 245 of Sugar. Ger, Jap and Russ, Rhubarb 10, Potassium Carbonate 1, Borax 1, Water 80, to 60 of filtrate add 20 of Cinnamon Water and 120 of Sugar. Swi— Rhubarb 10, Potassium Carbonate 1, Borax 1, Tincture of Cinnamon 12, Simple Syrup 176, Hung, Rhubarb 20, Sodium Carbonate 4, Diluted Spirit 20, Cold Water a sufficiency, to 200 of filtrate add 340 of Sugar. Ital (Scelopopo di Cicorione e Rabarbaro), Rhubarb 1, Juice of Chicory Leaves 12, Sugar 16, Swed, Rhubarb 5, Sodium Carbonate 1, Water a sufficiency, after filtration add to 57 of filtrate 63 of Sugar, Port (Carope de Rhubarbo), Rhubarb 2, Water 15, Sugar 60, Mex (Jirabe de Achicoria y Rhubarbo), Extract of Rhubarb 2, Simple Syrup 975, Belg, Fluid Extract of Rhubarb 50, Potassium Carbonate 5, Cinnamon Water 30, Simple Syrup 915, U.S., Fluid Extract 100, Spirit of Cinnamon 1, Potassium Carbonate 10, Water 50, Syrup to make 1000. All by weight except U.S.

Fr has a Compound Syrup and U.S. has also Syrupus Rhei Aromaticus.

TINCTURA RHEI COMPOSITA. COMPOUND TINCTURE OF RHUBARB

Rhubarb Root, 2, Cardamom Seeds, $\frac{1}{2}$, Coriander Fruit, $\frac{1}{2}$, Glycerin, 2, Alcohol (60 p.c.), q.s. to yield 20 (1 in 10)

B.P. 1880 contained Saffron, but no Glycerin

Dose — $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 cc for repeated administration, for a single administration, 2 to 4 fl drim = 7.1 to 14.2 cc

Foreign Pharmacopœias— Official in Austri (Tinctura Rhei Vinosa) and Hung (Tinctura Rhei Dorelli), Rhubarb 10, Orange Peel 2, Cardamom Seeds 1, Malaga Wine 100, in 100 of filtrate dissolve 15 of Sugar, Dutch (Vinum Rhei), Rhubarb 9, Cardamom 1, Malaga Wine 100, Ger (Tinctura Rhei Vinosa), Rhubarb 8, Orange Peel 2, Cardamom 1, Sherry 100 filter, and in the filtrate dissolve a seventh part of Sugar, Jap (Tinctura Rhei),

Rhubarb 10, Cassia 1, Cardamom 1, Alcohol 50, Distilled Water 50, Now and Swed (Tinctura Rhei Amara), Rhubarb 10, Gentian Root 4, Cardamom 1, Alcohol (64 p c), 100, Russ (Tinctura Rhei Vinosa), Rhubarb 8, Orange Peel 2, Cardamom 1, Sherry 100, Sugar 12, Swiss (Vinum Rhei Compositum), Rhubarb 8, Orange Peel 2, Cardamom 1, Vinum Meridianum Dulce 100, U S (Tinctura Rhei), Rhubarb 20, Cardamom 4, Glycerin 10, Alcohol and Water, each $q\ s$ to make 100, also (Tinctura Rhei Aromatica), Rhubarb 20, Saigon Cinnamon 4, Cloves 4, Nutmeg 2, Glycerin 10, Alcohol and Water, each $q\ s$ to make 100 All by weight except U S

Belg, Fl, Ital and Mex have a Simple Tincture, Rhubarb 1, Alcohol (60 p c) 5, Port, Rhubarb 1, Alcohol (65 p c) 5

See also Tinctura Rhei Aquosa, given under Infusum Rhei

Tests—Compound Tincture of Rhubarb has a sp gr of 0.970 to 0.975, it contains about 15 p c w/v of total solids and about 50 p c w/v of Absolute Alcohol

Not Official.

ELIXIR RHEI—Rhubarb Root, in No. 12 powder, 5, Fennel Fruit, bruised, 2, Glycerin 3, Refined Sugar 4, a mixture of Alcohol (90 p c) 1, and Distilled Water 3, $q\ s$ to produce 20—*B P C Formulary* 1901, now incorporated in the *B P C* with the *syn* Liquor Rhei Dulcis

Dose—1 to 3 fl drms = 3.6 to 10 G cc

EXTRACTUM RHEI COMPOSITUM—Ext Rhei 3, Ext Aloes 1, Resina Jalapæ $\frac{1}{2}$, Soap 2—*Ger*

Extract of Rhubarb 6, Extract of Aloes 2, Jalap Resin 1, Soap 1—*Austr, Dutch and Swiss*

Extract of Rhubarb 6, Extract of Barbados Aloes 2, Jalap Resin 1, Hard Soap 1—*B P C*

FLUIDEXTRACTUM RHEI.—100 of Rhubarb, in No. 30 powder, macerated in and subsequently percolated with a mixture of Alcohol (95 p c) 80, and Water 20, reserve the first 75 of the percolate, and evaporate the remainder to a soft extract, which mix with the reserved portion and make up with the menstruum to 100—*U S P*

Average Dose—15 minims = 0.9 cc

This has been incorporated in the *B P C*

INFUSUM RHEI CONCENTRATUM—Rhubarb, in No. 10 powder, 40, Alcohol (90 p c), 25, Dilute Chloroform Water (1 in 1000), $q\ s$ to make 100.—*Farr and Wright, P J '06, 1 165 and '07, 1 622, C D '06, 1 252, Y B P. 1907, 250* Prepare by repercolation

This appears in the *B P C*

MISTURA RHEI CUM SODA—Rhubarb Root, in powder, 5 grains, Sodium Bicarbonate 10 grains, Caraway Water to 1 fl. oz.—*St Thomas's*

This has been incorporated in the *B P C*

Mistura Rhei et Sodæ—Sodium Bicarbonate 3.5, Fluid Extract of Rhubarb 1.5, Fluid Extract of Ipecac 0.3, Glycerin 35, Spirit of Peppermint 3.5, Water $q\ s$ to make 100—*U S P*

PILULA RHEI ET COLOCYNTHIDIS ET HYDRARGYRI.—Compound Rhubarb Pill 1 grain, Compound Colocynth Pill 1 grain, Mercury Pill $\frac{1}{2}$ grain in each pill—*B P C*

PILULA RHEI ET NUCIS VOMICÆ—Compound Rhubarb Pill 3 grains, Extract of Nux Vomica $\frac{1}{2}$ grain, Alcoholic Extract of Belladonna $\frac{1}{2}$ grain in each pill—*St Thomas's*

The *B P C* use the same formula as above, the quantities being 2½ grains, $\frac{1}{2}$ grain, $\frac{1}{2}$ grain, respectively, with Milk Sugar to make a 4-grain pill

PULVIS RHEI CUM HYDRARGYRO—Rhubarb Root, in powder, 2 grains, Mercurous Chloride $\frac{1}{2}$ grain, Ginger, in powder, $\frac{1}{2}$ grain, dose for a child 12 months old—*St Thomas's*

This has been incorporated in the *B P C*

PULVIS HYDRARGYRI ET RHEI—Rhubarb, in powder, 3, Mercury with Chalk 1, Sodium Bicarbonate 3—*St Mary's*

Pulvis Rhei cum Hydrargyrio et Soda *Syn* Baid's Apertient Powder—Rhubarb Root 50, Mercury with Chalk 16 50, Sodium Bicarbonate *qs* to produce 100 Dose—6 to 12 grams—*L P C'*

PULVIS RHEI CUM SODA—Rhubarb Root, in powder, 1 gram, Sodium Bicarbonate 2 grams, dose for a child 12 months old—*St Thomas's*

This has been incorporated in the *B P C*

TINCTURA RHEI AQUOSA—Rhubarb 10, Potassium Carbonate 1, Sodium Borate 1, boiling Distilled Water 90, Alcohol 9, after the lapse of an hour strain the solution by applying a slight pressure, with every 85 parts of the strained liquid mix Cinnamon Water 15 Prepare freshly when required—*Ger* and *Jap*

This has been incorporated in the *L P C'*

Dutch Rhubarb 10, Borax 5, Spirits of Wine Diluted 20, Cold Water 80

Dutch, Rhubarb 10, Sodium Carbonate 2, Cinnamon Water to produce 100

Hung, Rhubarb 10, Sodium Carbonate 2, Cold Distilled Water 160, Alcohol (70 p.c.) 10

Russ, Rhubarb 10, Borax 1, Potassium Carbonate 1, Distilled Water Ebullient 85, Spirits of Wine (90 p.c.) 10, Cinnamon Water 15

Swiss, Liquid Extract 10, Borax 1, Potassium Carbonate 1, Alcohol 8, Cinnamon Water 20, Water 60

VINUM RHEI—Rhubarb Root in coarse powder, 1½ oz., Cancelli Bark 60 grams, Sherry 20 fl oz.—*L P C'* 1885, omitted in 1898

This has been incorporated in the *L P C'*, employing Detannated Sherry

Official in Belg, 1 of Fluid Extract in 20

PURGATIN, PURGATOL (Anthraquinum Diacetate) A yellow, or brownish yellow, micro-crystalline powder, insoluble in Water, sparingly soluble in Alcohol (90 p.c.) Introduced as a synthetic purgative belonging to the series of oxyanthraquinones

Useful in chronic constipation occurring along with neurasthenia, hypochondria, or hemorrhoids, where it is appropriately employed in place of Rhubarb and Aloe

Rumicin, dried extract from the Root of *Juncus crispus*, has been used as an eclectic preparation It has properties similar to Rhubarb, dose, 1 to 5 grains = 0.06 to 0.32 gramme

It must not be confounded with the crystalline substance Rumicin, which is allied to Chrysophanic Acid

RHCEADOS PETALA.

RED POPPY PETALS

Fr, COQUELICOT, Ger, KLAUSCHROSENBLUMEN, Ital, ROSOLACCIO, Span, AMAPOLA

The bright scarlet coloured, fresh Petals of *Papaver Rhæas*, L., possessing a peculiar narcotic odour and a mucilaginous bitter taste

Chiefly used as a colouring agent.

Official Preparation—Syrupus Rhæados

Foreign Pharmacopœias—Official in Aust, Belg, Dutch, Fr (Coque licot), Mex and Span (Amapola), Swiss

Descriptive Notes.—The fresh petals of *Papaver Rhæas* are Official There are several forms or allied species, all with red petals, but differing in the size of the flower and the shape and hairiness of the ovary, as well as in the shape of the leaf segments One of these

has a purplish-black spot at the base of each petal, but the scarlet colour of the petal is deeper than that of other forms, and this variety gives a deeper coloured syrup. The petals of the typical plant are crumpled when freshly unfolded, about $1\frac{1}{2}$ to 2 in (4 to 5 cm) broad, and of a bright red colour. They have a slightly bitter taste and characteristic odour.

Tests.—Red-Poppy Petals contain about 16 p c of ash.

Preparation

SYRUPUS RHILADOS. SYRUP OF RED-POPPY

Dissolve (with heat) 36 of Sugar in a strained infusion of Red-Poppy Petals, 13, in Distilled Water, 20, preserve with $2\frac{1}{2}$ of Alcohol (90 p c), total weight should be 58 (1 in 3 $\frac{1}{2}$)

Dose.— $\frac{1}{2}$ to 1 fl drim = 1.8 to 3.6 c c

This syrup is liable to fermentation, and is therefore preserved by the addition of 1 p c. In India and the Colonies the Alcohol may be increased up to twice the quantity ordered in the formula.

Foreign Pharmacopœias—Official in Dutch and Mex.

Not Official

RHUS TOXICODENDRON.

POISON IVY

The fresh Leaves of *Rhus radicans*, L., were official in *U S P* 1890, and a Tincture from them is given in doses of 1 to 5 minims for rheumatism. Fluid Extract (1 in 1) is also made.

Recently (August, 1908) attention has been called to the poisonous character of this plant, and that it produces an eruption in persons handling it, an alcohol solution of Lead Acetate applied to the rash affords speedy relief—*P J '08*, ii 232, 271.

Rhus glabra, L., (*U S*) and *Rhus aromatica*, Ait., have been used as tonics and astringents, given for nocturnal incontinence of urine. Both these can be supplied as Fluid Extracts (1 in 1), doses, 5 to 10 minims = 0.3 to 0.6 c c.

RICINI OLEUM.

CASTOR OIL

FR, HUILE DE RICIN, GER, RICINUSÖL, ITAL, OLIO DI RICINO, SPAN, ACEITE DE RICINO

A colourless, or pale yellow, almost odourless, thick viscid fluid, possessing at first a mild and subsequently somewhat nauseous taste; expressed from the Seeds of *Ricinus communis*, L.

It should be kept in well-closed vessels and protected as far as possible from exposure to the air, as it has a tendency to gradually thicken.

Reinoleic Acid is stated to be the active principle. The Seeds contain a toxic phytalbumose, Ricin, which is extremely poisonous, it is not contained in the Oil.

Solubility—Entirely soluble in all proportions of Absolute Alcohol, Ether, Oil of Turpentine and Glacial Acetic Acid, 1 in $3\frac{1}{2}$ of Alcohol (90 p c)

Medicinal Properties—A mild and speedy cathartic. It is the best purgative in constipation from indurated feces, or after swallowing toxic substances. Used in diseases attended with irritation or inflammation of the bowels, as colic, and diarrhoea due to indigestible food, dysentery and the constipation of typhoid fever, the most suitable purgative after parturition during pregnancy and after abdominal operations. The safest cathartic for infants, to whom a larger relative dose than to adults may be given, relieves infantile intestinal spasms. It may be administered in an enema with some mucilaginous or oily fluid.

Dropped into the eye it cures the irritation caused by a foreign body.

The decoction of the leaves of *Lacina* applied to the breast is said to produce an abundant secretion of milk.

Dose 1 to 8 fl. drms = 3.6 to 28.4 c c

Prescribing Notes—In draught suspended with mucilage of Gum Acacia, or in capsules (see below).

One of the best & most reliable mode of taking Castor Oil is to pour it on to some Milk or Cream contained in a wine glass, the interior and edges of which have been moistened with the latter.

It is used as a solvent for all kinds of bases in ophthalmic practice.

In the treatment of dysentery (*M. G. O.*, n. 219, 280), 1 oz. of the Oil, with 30 minims Tincture of Opium and Milk diet.

In infantile diarrhoea administered in small repeated doses. A formula which has been found convenient is Oleum Ricini 10 minims, Tincture of Rhubarb, 5 minims, Glycerum 5 minims, Tragus with $\frac{1}{2}$ gram Peppermint Water, to 1 to 1 drms to be given every 4 hours for the first 36 hours, and then less frequently. *Ph. Ann.* 1867.

Official Preparation—Mistura Olei Ricini. (Contained in Colloidium Flexile, Lammamentum Sinapis and Pilula Hydrargyri Subchloridi Composita.)

Not Official—Capule of Castor Oil, Emulsio Olei Ricini, Mistura Olei Ricini, Unguentum Olei Ricini, and Oleum Ricini Aromaticum.

Foreign Pharmacopœias—Official in all.

Tests—Castor Oil has a sp. gr. of 0.960 to 0.968. The *B. P.* states from 0.950 to 0.970. Good medicinal samples of the Oil never possess so low a gravity as 0.950. The *U. S. P.* gives the gravity of 0.941 to 0.961 at 25° C (77° F), the *P. G.* 0.950 to 0.970. Ten good medicinal samples of the Oil examined in the author's laboratory had a sp. gr. of 0.960 to 0.966 with an average of 0.963. When cooled to 0° C (32° F) a crystalline flocculent deposit settles out, and when reduced to a temperature of about -15° C (-9° F) it forms a yellowish buttery mass. When exposed to the air it gradually thickens and dries, forming a varnish. The Oil contains a certain proportion of free acid, which may be determined by dissolving a weighed quantity of the Oil (5 grammes) in Alcohol (90 p c) 25 c c, warming and titrating with Tenth normal Volumetric Sodium Hydroxide Solution. The 10 samples referred to above showed from 1.05 to 3.5 p c with an average of 2.1 p c. The Saponification value of the Oil ranges from 176 to 188. The *U. S. P.* gives 179 to 183, no figure is given in the *P. G.* The above-mentioned

10 samples showed from 176.4 to 187.6, with an average of 182.0. The Iodine absorption of the Oil varies from 85 to 90 p.c., the *U.S.P.* states not less than 84 nor more than 89, no figure is recorded in the *P.G.* The medicinal samples referred to above showed from 85.09 to 90.17, with an average of 87.5.

A determination of the optical rotation of the specimen affords a useful means of judging of its purity, the Oil is dextrogyrate, the optical rotation in a tube of 100 mm. being equal to $+4^{\circ}$ to $+4.5^{\circ}$. The *B.P.* does not make any mention of a determination of free acid, it gives no figures for the Saponification value or for the Iodine absorption, nor does it refer to the optical rotation.

The more general impurities are fixed Oils other than Castor, such as Cottonseed Oil, Lard Oil, etc. Castor Oil is an exception to the usual characters of the fixed Oils in regard to its solubility in Alcohol. It should dissolve completely in all proportions of Absolute Alcohol, in Glacial Acetic Acid, and in $3\frac{1}{2}$ times its volume of Alcohol (90 p.c.), indicating the absence of more than about 5 p.c. of fixed Oils other than Castor. The *B.P.* gives a test with Sulphuric Acid for detecting the presence of various fixed Oils including Cottonseed, and requires that when 3 c.c. of the Oil are dissolved in 3 c.c. of Carbon Bisulphide, the mixture should not assume a brown colour when shaken with 1 c.c. of Sulphuric Acid; the test is generally found to be quite unreliable. A useful test for detecting the presence of other fixed Oils is that with Petroleum Ether, but not when carried out as directed in the *B.P.* The latter states that equal volumes of Castor Oil and Petroleum Ether do not yield a clear mixture if kept at 15.5°C (60°F), but that they yield a perfectly clear mixture if other fixed Oils be present. The *U.S.P.* description of the test is the more correct, it states that, when mixed with an equal volume of Petroleum Benzine the Oil yields at 17°C (62.6°F) a clear solution, but that at 15°C (59°F) it forms a turbid mixture. It has been remarked that the monograph requires complete revision, Saponification and Iodine values should be introduced, the Sulphuric Acid test needs revision, if retained, but is of little service.

Carbon Bisulphide and Sulphuric Acid—If 3 c.c. of the Oil be shaken with 3 c.c. of Carbon Bisulphide and 1 c.c. of Sulphuric Acid, the mixture should not acquire a brown colour, *B.P.*, a blackish-brown colour, *P.G.* and *U.S.P.*

Iodine Absorption—If 0.3 gramme of the Oil be dissolved in 10 c.c. of Chloroform in a 250 c.c. bottle or flask and 25 c.c. of a mixture of equal volumes of Alcoholic Iodine T.S. and Alcoholic Mercuric Chloride T.S. added, and if after standing for 9 hours protected from light, 20 c.c. of Potassium Iodide T.S. be introduced and the mixture diluted with 50 c.c. of Water, on titrating the excess of Iodine with Tenth-normal Sodium Thiosulphate Volumetric Solution an Iodine value of not less than 84 nor more than 89 should be obtained, *U.S.P.*

Preparation.

MISTURA OLEI RICINI. CASTOR OIL MIXTURE.

To $1\frac{1}{2}$ of Mucilage of Gum Acacia add with trituration in small portions alternately, 3 of Castor Oil and a mixture of Orange-Flower Water (undiluted) 1 and Cinnamon Water $2\frac{1}{2}$.

The Oil is now emulsified by means of Mucilage of Gum Acacia in place of saponification with Solution of Potassium Hydroxide, and Cinnamon Water replaces the Oils of Lemon and Cloves

Dose Is a draught, 1 to 2 fl oz = 28.4 to 56.8 cc

Not Official

CAPSULES OF CASTOR OIL —Flexible capsules containing 30 minims, or 60 minims in each

EMULSIO OLEI RICINI —Castor Oil 3 fl oz, Mucilage of Acacia, 3 fl oz, Syrup of Ginger, 1 fl oz, Cinnamon Water, 1 fl oz —*Squire*
Castor Oil 1 fl oz, Yolk of Egg, 1 fl oz, Syrup, 1 fl oz, Peppermint Water 1 fl oz —*Squire*

Either of the formulae yield a good emulsion

MISTURA OLEI RICINI *Syn* Emulsio Olei Ricini —Castor Oil, 6 fl drim, Mucilage of Gum Acacia, 3 fl drim, Orange Flower Water, 2 fl drim, Cinnamon Water, to make 2 fl oz —*B.P.C.*

ENEMA OLEI RICINI —Castor Oil, 2 fl oz, Mucilage of Starch, 18 fl
Castor Oil 1 fl oz, Olive Oil, 5 fl oz

OLEUM RICINI AROMATICUM —Geluside 7½ grains, Sodium Bicarbonate, 7½ grains, Chloroform 10 minims, Oil of Pimento, 75 minims, Oil of Cassia 75 minims, Oil of Clove, 75 minims, Castor Oil, *qs* to make 10 fl oz —*Canadian Formulary*

Amyl Acetate 0.1, Geluside 0.15, Alcohol (90 per cent), 5, Castor Oil, *qs* to produce 100 —*L.P.C.*

ROSÆ GALLICÆ PETALA.

RED ROSE PETALS

FR, ROSE ROUGE, GER, ROSEHÜHN, ITAL, ROSA ROSSA, SIAM, ROSA ROJA

Dark purplish red, velvety, claw shaped petals, possessing a roseaceous odour, and a slightly acidulous, bitter, astringent taste. They usually occur in small, crumpled, conical masses, and are officially described as the fresh and dried unexpanded Petals of *Rosa Gallica*, L., from cultivated plants

Medicinal Properties.—Used on account of their colouring matter and mild astringency

Prescribing Notes The Acid Infusum is prescribed with Glycerin of Tannin or Alum as an astringent vehicle, it also forms a suitable vehicle for Magnesium Sulphate, the Syrup is used as a colouring agent, and the Confection as a pill excipient. The Astringent Infusum is given with Quinine

Official Preparations Of the petals, Confectio Rosæ Gallicæ, Infusum Rosæ Acidum, and Syrupus Rosæ. The confection is contained in Pilula Aloes Barbadosens, Pilula Aloes et Asafetide, Pilula Aloes Socotrine, and Pilula Hydragryi

Not Official Fluidextractum Rosæ, Infusum Rosæ cum Acido Nitrico, Infusum Rosæ Acidum Concentratum, Mel Rosæ, Pulvis Rosæ Compositus, and Unguentum Rosatum

Foreign Pharmacopœias —Official in Austr, Belg, Dutch, Fr, Ger, Hung, Ital, Jap, Port, Russ, Span, Swiss and U.S.

Descriptive Notes —There are several varieties of *Rosa Gallica* in cultivation, the flowers of which are met with in commerce. The petals of those cultivated in England obtain a higher price than 'Exotic

petals, &c., those imported from France, Germany, Holland, etc., having a brighter red colour, a greater fragrance and being less broken. The dried petals consist of the flower bud with the lower or calycine portion removed, and the buds are more or less broken up in drying, they have a purplish-rose tint, but are yellowish towards the base. The taste is feebly acid, astringent, and slightly bitter. The fragrance depends upon the variety under cultivation, that known as General Jacqueminot affording a fragrant product of good colour, but the colour depends partly also upon the maturity of the petal and the care taken in drying. Interior specimens coloured with aniline dyes are sometimes offered, but these are readily detected by the absence of the yellow base of the petal, as the whole becomes reddened by the dye.

Tests.—Red-Rose Petals leave about 4 p.c. of ash when incinerated with free access of air.

Preparations

CONFECTIO ROSÆ GALLICÆ. CONFECTION OF ROSES

Fresh Red-Rose Petals, 1, Refined Sugar, 3 (1 in 4)

Used as a pill basis. Also applied in aphthous conditions of the mouth.

Official in U.S.

INFUSUM ROSÆ ACIDUM. ACID INFUSION OF ROSES

Red-Rose Petals, dried and broken, $\frac{1}{2}$ oz., Diluted Sulphuric Acid, 2 fl. drms., Distilled Water, boiling, 20 fl. oz., infuse 15 minutes (1 in 40)

A similar infusion was in use in 1674.

Prescribed with Alum it forms a good gargle, but Borax or Alkalis change the colour to green.

Dose.— $\frac{1}{2}$ to 1 fl. oz. = 7.1 to 14.2 cc.

Foreign Pharmacopœias.—Official in Port (Infusão de Rosas Compostas), Red-Rose Petals, 5, Diluted Sulphuric Acid, 2, Boiling Water, 200.

SYRUPUS ROSÆ. SYRUP OF ROSES

Dissolve (with heat) 30 of Sugar in an infusion of dried Red-Rose Petals, 2, Refined Sugar, 80, boiling Distilled Water, 20, the total weight should be nearly 46 (1 in 17 $\frac{1}{4}$).

Dose.— $\frac{1}{2}$ to 1 fl. drms. = 1.8 to 3.6 cc.

Foreign Pharmacopœias.—Official in Belg., Fluid Extract 1, Simple Syrup 9. Mex. made from Rosa Centifolia, U.S., Fluid Extract 125, Dilute Sulphuric Acid 10, Sugar 750, Water, q.s. to make 1000.

Not Official.

FLUIDEXTRACTUM ROSÆ.—1000 grammes of Roses, in No. 20 powder, percolated with a mixture of 100 cc. Glycerin, and 900 cc. of Diluted Alcohol (Alcohol 50 p.c.) until the powder is exhausted. Reserve the first 750 cc., and evaporate the remainder, at a temperature not exceeding 50° C. (122° F.), to a thick extract. Dissolve this in the reserved portion, and make up with Diluted Alcohol to 1000 cc. —U.S.P.

Official in Belg.

This has been incorporated in the B.P.C., employing Alcohol (50 p.c.)

INFUSUM ROSÆ ACIDUM CONCENTRATUM.—Dried Red-Rose Petals, in No. 20 powder, 20, Diluted Sulphuric Acid and Alcohol (20 p.c.), of

each sufficient to make 100 Moisten the powder with some of the Alcohol containing one fortieth its volume of Diluted Sulphuric Acid, macerate for 2 hours, then pack in a glass percolator and percolate slowly with more of the Acidulated Alcohol until 92½ has been collected Add to this 7½ of Diluted Sulphuric Acid, set aside for 7 days, filter Dose — ½ to 1 fl drm = 18 to 36 cc — *Pharm* and *Wright, P J* '06, 1 165, and '07, 1 622, *C D* '06, 1 252, *Y B P* 1907, 251

This appears in the *B P C*

INFUSUM ROSÆ CUM ACIDO NITRICO — Rose Petals, broken small, 2, Diluted Nitric Acid, ½, cold Distilled Water, 40, infuse 2 hours, frequently stirring, strain, and add Powdered Sugar, 1

MEL ROSÆ — Fluid Extract of Roses 12 cc, Clarified Honey, a sufficiency to make the product weigh 100 grammes — *U S P*

This has been incorporated in the *B P C*

Foreign Pharmacopœias — Official in Ger and Jap, 1 of Rose Leaves is macerated with 5 of Alcohol (90 p c) for 24 hours, express and filter, mix with the nitrate 9 of Purified Honey and 1 of Glycerin, and evaporate to 10, both by weight Mel Rosatum is also official in Austr, Dutch, Fr, Mex and Swiss, but the formulas differ a good deal from one another

PULVIS ROSÆ COMPOSITUS — Oil of Rose and Chloroform, of each 1 (or combined 4 drops), Acacia, 145 grains, Sugar, 840 grains, Solution of Carmine, 13 drops Useful as an agreeable diluent for powders such as Calomel, Grey Powder, and Jalapin, also as a colouring and flavouring agent in mixtures, ¼ or ½ oz in 6 oz — *Martindale*

This has been incorporated in the *B P C* as follows — Oil of Rose, 0 10; Gum Acacia, in powder, 15, Solution of Carmine, 1 25, Refined Sugar, in powder, *q s* to produce 100 — *B P C*

UNGUENTUM ROSATUM — Alkanet Root, crushed, 13 grains, Otto of Roses, 1 minim, White Wax, 4 grains, Prepared Lard, 1 oz

Alkanna Root, bruised, 3, White Beeswax, 1, Oil of Rose, 0 20, Lard, *q s* to produce 100 — *B P C*

ROSÆ OLEUM.

OIL OF ROSE

B P Syn — OTTO OF ROSE

FR, ESSENCE DE ROSL, GER, ROSINOL, ITAL, ESSIENZA DI ROSA,
SPAN, ESENCIA DL ROSA

At a temperature of about 30° C (86° F), it is a pale yellow, or greenish-yellow, oily liquid, of about the consistency of Almond Oil It has a very powerful rosaceous odour and somewhat sharp taste At temperatures between 18° to 21° C (64 4° to 69 8° F), shining, acicular crystals, or glistening crystalline laminae, separate out, and when further cooled the Oil sets to a semi-solid crystalline mass, which again melts when gently warmed

It is officially described as the Oil distilled from the fresh plant of *Rosa damascena*, Miller The *U S P* describes it as a volatile Oil distilled from the fresh flowers of *Rosa damascena*, Mueller, and requires it to possess, when assayed by the process described in small type below, a Saponification value of not less than 10 nor more than 17 The *P G* describes it as a volatile Oil from the corolla of some varieties of roses, without defining any species

It should be kept in well-stoppered glass bottles of a dark amber tint in a cool place and protected as far as possible from the light

The Oil should be completely liquefied by heat and well mixed before being used for dispensing purposes

The average composition of Otto of Rose is stated to be Geraniol 40 p c, Citronellol 28 p c, Phenyl-ethyl Alcohol, 1 p c, Stearoptene 18 to 19 p c and small quantities of Linalool, Citral, Normal Nonylic Aldehyde, and other bodies

The vehicle of the odour is the elæoptene (Rhodinol) alone, and the less stearoptene there is in an otto used for manufacturing purposes the better — *C D '96*, 11 349

Medicinal Properties.—The principal use in pharmacy is as a perfume in various preparations

Official Preparation—Contained in Unguentum Aquæ Rosæ

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Fr, Ger, Hung, Jap, Mex, Port, Russ, Swiss and U S

Tests—Rose Oil is officially required to possess a sp gr. of 0 856 to 0 860 at 30° C (86° F), the statement referring to the sp gr of the Oil at 30° C (86° F) as compared with Water at 15 5° C (60° F), *vide Digest of Researches and Criticisms Report for 1898* The *USP* gives a sp gr of 0 855 to 0 865 at 25° C (77° F), the *PG* gives no figures for the sp gr The *Fr Codex* (1908) gives 0 855 to 0 865 at 20° C (68° F) The *BP* limit of gravity is generally considered too high, it usually falls from 0 850 to 0 858 The Oil is laevogyrate, the optical rotation of good specimens being from -1° 30' to -3°, neither the *USP* nor the *PG* includes a determination of the optical rotation The congealing point lies between 19° and 22° C (66 2° and 71 6° F) The *BP* states that the congealing and melting points vary, the proportion of crystalline matter, but should lie between 18° and 22 2° C (67° and 72° F), the *USP* gives the congealing point as between 18° and 22° C (64 4° and 71 6° F), and gives specific instructions as to the method to be adopted in determining the congealing point, which instructions appear below The *PG* states that crystals commence to separate out at 18° to 21° C (64 4° to 69 8° F), melting again at a higher temperature The *Fr Codex* (1908) gives 23 5° C (74 3° F) The refractive index of the Oil lies between 1 459 and 1 464, neither the *BP*, *USP*, nor *PG* refers to the refractive index Useful information is afforded of the nature of an Oil by a determination of the Acid and Ester value, the Acid value varies from 0 5 to 3 and the Ester value from 8 to 16, the *BP* makes no reference either to the Acid or Ester value The *USP* does not include an Ester value, but requires the Saponification value to be not less than 10 nor more than 17 as determined by the process given in small type below The *PG* does not include either an Acid or an Ester value The Oil contains from 18 to 23 p c of Stearoptene, and when the Stearoptene is carefully separated and purified it possesses a m p of from 33° to 35° C (91 4° to 95° F) The Iodine absorption has been suggested (*Analyst* '04, 175, *C D '04*, 1 398, '04, 11 703) as a means of detecting adulterated samples The Iodine absorption of genuine Otto was found to range from 187 to

194, that of artificial Oil from 221 to 254 for Oils containing Stearoptene, for those without Stearoptene, 261 to 279 Further information concerning the genuineness or otherwise of a specimen may be obtained by a determination of the percentage of Geraniol or Citronellol by acetylsation The percentage of Geraniol generally present in genuine samples varies from 65 to 75 p c, and occasionally may be as high as 76 p c Citronellol ranges from 25 to 35 p c, Geraniol from 30 to 33 p c

The chief and most commonly occurring adulterant of Otto of Rose is Turkish Geranium Oil, the presence of which may be determined by the alteration which it causes in one or more of the above constants of the Oil Geranium Oil lowers the sp gr and increases the Ester value, it also lowers the congealing point In addition to Geranium Oil a specimen may contain Spermaceti, Paraffin Wax, Palma Rosa Oil and Guaiacum Wood Oil Spermaceti and Paraffin Wax may be detected by a determination of the m p of the Stearoptene, and also the determination of its amount Spermaceti, if present, may be recognised by a determination of the Saponification value of the separated Stearoptene, Spermaceti absorbing an appreciable amount of Potassium Hydroxide on saponification Palma Rosa Oil, if present, may be detected by its influence on the Saponification value, and also by its effect on the Alcohol-content as determined by acetylsation Guaiacum Wood Oil may be detected by the microscopical appearance of the crystals separating from the Oil on cooling, and by the isolation and a determination of the m p of the Alcohol, Guaiol, the pure Alcohol melts at 91° C (195 8° F) Guaiacum Wood Oil tends to increase the sp gr and the optical rotation of the Oil, to raise the congealing point and to slightly lower the Saponification value Guaiacum Wood Oil leaves on evaporation a resinous mass amounting to about 16 2 p c Its presence is also indicated by the m p and by a determination of the Acetyl value of the Stearoptene

The Oil of White Rose is stated to contain a large percentage of Stearoptene, and has therefore been used to rectify the decrease in Stearoptene-content caused by the addition of other adulterants Parry does not see that a White Rose product should be regarded as an adulteration because it yields a few per cent more Stearoptene The *BP* monograph requires complete revision

Determination of Melting Point—Introduce about 10 c c of Oil into a test tube of about 15 mm diameter, insert a thermometer in such a manner that it touches neither the bottom nor the sides of the tube Raise the temperature of the Oil in the tube from 4° to 5° above the saturation point by grasping it in the hand, and shake the tube gently Allow the Oil to cool, and, when the first crystals appear, note the temperature This is regarded as the congealing point, a second test should be made for confirmation, *U S P*

Volumetric Determination—A measured quantity of 2 c c of the Oil is accurately weighed out in a weighing bottle and transferred by means of a little Alcohol (94 9 p c) to a flask having a capacity of about 100 c c, 20 c c of Semi-Normal Volumetric Alcoholic Potassium Hydroxide Solution added, and after connecting with a reflux condenser, the mixture boiled during half an hour on a water bath, the mixture is then cooled, diluted with 50 c c of Distilled Water, a few drops of Phenolphthalein TS added, and the excess of Volumetric

Alkali Solution titrated with Semi-Normal Volumetric Sulphuric Acid Solution
The number of cc of Semi-normal Volumetric Sulphuric Acid Solution is subtracted from 20, the difference is multiplied by 27.87, and the product divided by the weight of Oil taken, the result being the Saponification value of the Oil—*U S P*

ROSÆ AQUA.

ROSE WATER

FR, EAU DISTILLÉE DE ROSE, GER, ROSEWASSER, ITAL, ACQUA
DISTILLATA DI ROSE, SPAN, AGUA DESTILADA DE ROSAS

A clear, colourless liquid, possessing a strong rosaceous odour, prepared by distillation from the flowers of *Rosa damascena*, Miller, and diluted, immediately before use, 1 to 2 of Distilled Water

The Rose Water of commerce is a saturated solution of the essential Oil of the Rose flowers

Medicinal Properties.—An agreeable vehicle for medicines, employed in making lotions and eye-washes

Official Preparation.—Unguentum Aquæ Rosæ Contained in Mistura Ferri Composita, and the 'Rose Basis' for Lozenges

Foreign Pharmacopœias.—Official in Austr, Oil 5 drops, Warm Water 1000 grammes, Belg, Oil 0.8, Warm Water 1000, Dan, Oil 1, Tepid Distilled Water 10,000, Dutch, Oil 1, Water 5000, Fr, Mex, Port and Span, 1 of petals in 1, Ger and Jap, Oil 4 drops, Tepid Distilled Water 1000 cc, Ital, 1 in 2, Swiss, the Rose Water of commerce, undiluted, U S (Aqua Rosæ Fortior), the Rose Water of commerce (Aqua Rosæ), diluted with equal parts of Water

Preparation

UNGUENTUM AQUÆ ROSÆ. ROSE WATER OINTMENT
NO Syn—COLD CREAM

Heat until dissolved, Beeswax $1\frac{1}{2}$ oz, Spermaceti $1\frac{1}{2}$ oz, and Almond Oil (by weight) 9 oz, transfer to a warmed mortar, and add gradually with trituration Rose Water (undiluted) 7 fl oz, finally mix in 8 minims of Oil of Rose, and continue stirring until cold

A similar formula occurs in several of the Foreign Pharmacopœias, see p 357

Foreign Pharmacopœias.—Official in Mex, Rose Petals 1, Hog's Fat 1, Span, digested with an equal weight of Hog's Fat at a gentle heat for 3 days U S, White Wax 120, Expressed Oil of Almond 560, Codum Borate Water 190, Fr, (Cérat de Galien), White Wax 10 Almond at 25, all by weight also Cérat à la Rose, White Wax Vaseline 100 grammes, Caramine 1 gramme, Vaseline Oil 4 grammes, 20 drops

ROSMARINI OLEUM.

OIL OF ROSEMARY

NO Syn—OLEUM ANTHOS

FR, ESSENCE DE ROMARIN GER, ROSMARINOL, ITAL, ESSENZA DI
ROSMARINO, SPAN, ESENCIA DE ROMERO

A colourless, pale yellow, oily, limpid liquid, possessing a characteristic camphoraceous odour, and an aromatic and cooling taste. It is distilled from the Flowering Tops of *Rosmarinus officinalis*, L

Neither the *BP* nor the *PG* requires the Oil to contain any definite amount of Ester or of total Borneol. The *USP* requires it to contain not less than 2.5 p.c. of Ester calculated as Borneyl Acetate and not less than 10 p.c. of total Borneol.

It should be kept in well-stoppered glass bottles of a dark amber tint in a cool place and protected as far as possible from the light.

Rosemary Oil contains from 5 to 6 p.c. of Esters, chiefly Borneyl Acetate, and from 15 to 20 p.c. of Borneol. It also contains a mixture of dextro and lævo-Pinene, Camphene, Cineol and Camphor. The *BPC* states that the chief constituents are about 6 p.c. of Borneol and from 17 to 20 p.c. of Borneyl Acetate and other esters.

That distilled in Britain is superior to the imported.

Solubility—In all proportions of Absolute Alcohol, 2 in 1 of Alcohol (90 p.c.), sparingly in Alcohol (60 p.c.)

Medicinal Properties—Aromatic and carminative. It is used in hair lotions and liniments as a stimulant, also used for its odour, which is disliked by insects.

Dose— $\frac{1}{2}$ to 3 minims = 0.03 to 0.18 c.c.

Official Preparations—*Spiritus Rosmarini*. Contained in *Linummentum Saponis* and *Tinctura Lavandulae Composita*.

Foreign Pharmacopœias—Official in Austr., Dutch, Ger., Hung., Jap., Russ., Swiss and U.S. (*Oleum Rosmarini*), Belg. (*Rosmarini Essentia*), Dan., Norw. and Swed. (*Aether-oleum Rosmarini*), Fr. (*Essence de Romarin*), Ital. (*Essenza di Rosmarino*), Port. (*Essencia de Alecrim*), Span. (*Essencia de Romero*). Not in Mex.

Tests—Rosemary Oil has a sp. gr. of 0.900 to 0.920. The *BP* states 0.900 to 0.915, but the latter figure is regarded as too stringent. The *USP* states 0.894 to 0.912 at 25° C (77° F), the *PG* not under 0.900. It is dextrogyrate, the optical rotation being from +1° to +18°. The *BP* gives the optical rotation as not more than +10° in a tube 100 mm. long. The *USP* states that the angle of rotation shall not be more than +15° in a 100 mm. tube at a temperature of 25° C (77° F), *PG* does not give a figure for the optical rotation of the Oil. It is soluble in all proportions of Absolute Alcohol, and should dissolve in twice its volume of Alcohol (90 p.c.). The *USP* states that it is soluble in about one half volume or more of Alcohol (90 p.c.), also in 2 to 10 volumes of Alcohol (80 p.c.). The *PG* states that the Oil should afford a clear solution in half its weight of Alcohol (90 p.c.). Neither the *BP* nor the *PG* gives a method for determining the proportion of Ester in terms of Borneyl Acetate nor the total Borneol present, the former may be determined by saponification, the latter by acetylation. The *USP* employs the saponification and acetylation process described in small type below.

The more generally occurring impurities are Turpentine Oil, Petroleum Oil and Alcohol. Turpentine, if present in considerable quantity, may be detected by the optical rotation of the sample and if present in small proportion by the optical rotation of the first 10 p.c. of the distillate. It also causes a diminution in the sp. gr.

French Turpentine Oil is indicated by the Oil assuming a lævo-rotation, or by the lævo-rotation of the first 10 p c yielded on distillation. The *USP* requires the first 10 p c fraction to be dextrogyrate. Petroleum Oil is detected by the diminished solubility of the Oil in Alcohol (90 p c), and by the residue left on evaporating the Oil on a water-bath. Pure Rosemary Oil leaves only a slight amount of residue of a resinous character. The presence of Alcohol may be detected by the addition of a little solid Magenta. Magenta imparts no colour to the pure Oil, but the dye dissolves in the presence of Alcohol. Fractions of Camphor Oil have also been met with as adulterants of Oil of Rosemary, but their presence may be detected by the influence on the rotatory power or their sp gr, or their effect on the solubility of the Oil in Alcohol (90 p c).

Notwithstanding the official requirements, as well as those of other Pharmacopœias, that the Oil should be dextrogyrate, undoubtedly genuine samples are found which are lævogyrate.

According to Parry (*CD* '06, 1 671) the lævorotatory constituent occurs in greater proportion when the stalks are included, and an inferior Oil is then obtained. Oils derived from carefully picked leaves yield fractions which are lævogyrate, the genuine lævorotatory Oil, containing a comparatively low percentage of Borneol, may be assumed to have been distilled from both leaves and stalks. A dextro-rotatory Oil may, moreover, yield lævorotatory fractions, a first fraction of 10 p c having a lævo-rotatory optical rotation. The Spanish Pharmacopœia states that the Oil is lævogyrate. Schimmel is of opinion that in any case it will be well to continue exercising care in dealing with lævorotatory Rosemary Oils. He reports an authenticated sample of English Oil, examined in their laboratory, which possessed a sp gr of 0.9042, an optical rotation of $-2^{\circ} 49'$, an Ester value of 9.7, and which was soluble 1 in about 5 of Alcohol, (80 p c), with very slight turbidity, the optical rotation of the first 10 p c distillate was $-6^{\circ} 10'$. Samples of English Oils distilled during the years 1905, 1906, 1907 were indisputably genuine in character, and possessed optical rotations of $-0^{\circ} 24'$ to $-2^{\circ} 48'$.

Specimens of the various imported varieties examined in the author's laboratory in June 1893 showed optical rotations as follows —

Eperte	price 3s 1d	per lb, rotation	-8°	soluble in S V R	2 in 1
Extra	" 2s 6d	" "	-12°	" " "	2 in 1
Super	" 1s 9d	" "	-33°	" " "	2 in 9
Fine	" 1s 3d	" "	-40°	" " "	2 in 10
French Turpentine		" "	-57°	" " "	2 in 8

Specimens examined in the author's laboratory within recent years all possessed a dextro-rotation varying from $+7^{\circ}$ to $+11^{\circ}$.

Volumetric Determination of Esters—A measured quantity of 10 c c of the Oil is introduced into a tared flask, and its weight accurately determined. A measured quantity of 25 c c of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution is added, the flask connected with a reflux condenser, and the mixture boiled for 1 hour. It is then allowed to cool, and the excess of Semi-

normal Volumetric Alcoholic Alkali Solution is titrated with Semi normal Volumetric Sulphuric Acid, using Phenolphthalein T S as an indicator of neutrality. The number of c c of Semi normal Volumetric Sulphuric Acid Solution required is subtracted from 25, the difference multiplied by 9.734, and the product divided by the weight of Oil taken, the quotient representing the percentage of esters present in the Oil, expressed in terms of Borneol Acetate. The residual Oil from the saponification, is washed repeatedly with Water, transferred to an acetylation flask, mixed with 10 c c of Acetic Acid Anhydride and about 1 gramme of anhydrous Sodium Acetate, and boiled gently for 1 hour. The mixture is allowed to cool, the acetylated Oil is washed with Distilled Water and subsequently with Sodium Hydroxide T S until it is slightly alkaline to Phenolphthalein T S, and is then dried by means of fused Calcium Chloride and filtered, *U S P*

Volumetric Determination of total Borneol—A measured quantity of 5 c c of the dry acetylated Oil prepared as above is transferred to a tared flask, of about a capacity of 100 c c, and the weight accurately determined. 50 c c of Semi normal Volumetric Alcoholic Potassium Hydroxide Solution is added, the flask connected with a reflux condenser, the mixture boiled for 1 hour, when cooled the excess of Semi normal Volumetric Alcoholic Alkali Solution is titrated with Semi normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. The number of c c of Semi normal Volumetric Sulphuric Acid Solution required is subtracted from the number of c c of Semi normal Volumetric Alcoholic Potassium Hydroxide used (50 c c). The difference is multiplied by 7.649, and the product divided by the weight of dry acetylated oil employed, less the product of the multiplication of the number of c c of Semi normal Volumetric Alcoholic Potassium Hydroxide Solution absorbed by the acetylated oil by 0.021, the quotient represents the total percentage of Borneol present in the specimen under examination, *U S P*

Preparation

SPIRITUS ROSMARINI SPIRIT OF ROSEMARY

Oil of Rosemary, 1, Alcohol (90 p c), *q s* to yield 10 (1 in 10)

In *B P* '85 it was 1 in 50

Dose—5 to 30 minims = 0.3 to 1.8 c c

Foreign Pharmacopœias—Official in Austr, from leaves, Jap, 1 in 9, Port (Espirito d'Alecrim), Rosemary 5, Water 2, Alcohol (85 p c) 10, Mex (Alcoholato de Romero compuesto), Dried leaves 1, Lavender 1, Alcohol (80 p c) 10, Water 2, Russ, 1 in 100, Span (Alcohol de Romero), Rosemary 1, Alcohol (60 p c) 2, Swiss (Spiritus Rosmarini Compositus), Lavender 1, Peppermint 1, Rosemary 1, Salvia 1, Wormwood 1, Alcohol 20, Water 50

Unguentum Rosmarini Compositum official in Ger, Lard 16, Mutton Fat 8, Yellow Wax 2, Expressed Oil of Nutmeg 2, Oil of Rosemary 1, Juniper Oil 1, Swiss, Oil of Rosemary 1, Oil of Turpentine 3, Juniper Oil 6, Oil of Laurel 10, Yellow Wax 24, Lard 56

Not Official
RUTA OLEUM

OIL OF RUD

Fr, ESSENCE DE RUE, GER, RAUTENOL, ITAL, ESSENZA DI RUTA, SPAN, AÇEITE DE RUDA

A colourless, or pale yellow, oily liquid, possessing an intense, persistent, characteristic odour. It is distilled from the fresh Herb of *Ruta graveolens*, L.

Medicinal Properties—Antispasmodic. A topical stimulant and rubefacient. Administered in the form of enema for flatulent colic in children.

Dose,—1 to 4 minims = 0.06 to 0.24 c c,

Foreign Pharmacopœias—Official in Belg, Port and Span

French Oil of Rue is stated by Schimmel to differ from the Algerian Oil in that both contain about 90 p c of ketones, but the French Oil exclusively Methyl-nonyl-ketone, m p 15° C (59° F), the Algerian, Methyl heptyl-ketone m p -16° C (3° F)

Power and Lees in the examination of an essential Oil of Rue, apparently of Algerian origin, found the following constituents—Methyl-*n* heptyl ketone, Methyl-*n*-nonyl ketone, Methyl-*n* heptylcarbinol, Methyl-*n*-nonylcarbinol, a blue Oil, b p 180° C, boiling point, Acetic Acid in combination with Alcohols, an odour of Quinine, a mixture of free fatty acids, Methyl Salicylate, an ester of Valerianic Acid, Methyl Valerianate, Pinene, Lavandulonic and Cineol. The two alcohols represented about 80 p c of the Oil, and were present in about equal amounts. The two alcohols represented about 10 p c and were present partly in the uncombined state and partly as Acetic esters, the Methyl heptylcarbinol predominating. The two Terpenes, together with Cineol represented about 1 p c of the Oil. There was very little Pinene, and the amounts of Limonene and Cineol were about equal. The amount of blue Oil was about $\frac{1}{2}$ p c, and finally there was separated from the non-ketonic portion of the Oil a small amount of undistillable viscous substance, which was probably a decomposition product.—*Report of the Wellcome Chemical Research Laboratories*

Tests—Oil of Rue has a sp gr of 0.833 to 0.840, it is dextrogyrate possessing an optical rotation in a 100 mm tube of $+2^{\circ}$, the solidifying point is 8° to 10° C (46° to 50° F). It dissolves to form a clear solution in an equal volume of Alcohol (90 p c) or in 2 to 3 parts of Alcohol (70 p c). It was official in the B P 85.

CONFECTIO RUTÆ.—Fresh Rue, bruised, $1\frac{1}{2}$ oz, Caraway Seeds, $1\frac{1}{2}$ oz, Bay Berries, $1\frac{1}{2}$ oz. Prepared Sagipenum $\frac{1}{2}$ oz, Black Pepper, 2 drms, Honey, 16 oz. Distilled Water, as much as may be necessary.—P L 1851

This has been incorporated in the B P C

ENEMA RUTÆ—Confection of Rue, 3 drms, Infusion of Chamomile, to make 20 fl oz.—*St George's*

Confection of Rue, 1, Decoction of Bailey, q s to produce 100.—B P C

Oil of Rue, 30 minims, Starch Enema, 6 oz.—*Westminster*

Not Official**SABINÆ CACUMINA****SAVIN TOPS**

FR, SABINE, GER, SÄDERBAUMSPITZEN, ITAL, SABINA, SPAN, SABINA

The fresh and dried Tops of *Juniperus Sabina*, collected in spring from plants cultivated in Britain. The Savin Tops imported from France are not always those of *J. Sabina*.

It was official in B P '85

Medicinal Properties—A powerful local and general irritant. The ointment is used for maintaining discharges from granulating or blistered surfaces. It is a powerful emmenagogue, but its use requires caution, as it may cause inflammation of the abdominal and pelvic viscera.

Dose—4 to 10 grains = 0.26 to 0.65 grammes

Antidotes—Stomach-tube, emetics, Castor Oil, Linseed poultices to the abdomen, opiates and demulcents

Foreign Pharmacopœias—Official in all except Ger, Jap, Span and Swed

Descriptive Notes—Savin Tops are generally imported in this country in the fresh state from the cultivated shrub, but in some cases exist two or three varieties, less frequently in the dried state, in the form of woody twigs or branchlets, 6 to 9 inches (15 to 22.5 cm) long or more or imported from Italy

and the south of France in the form of broken twigs freed from the woody portion. In the Savin cultivated in England the leaves are generally spreading, but in the exotic Savin they are closely imbricated so as to form nearly cylindrical branchlets. The leaves are narrowly triangular and concavo-convex, about $\frac{1}{2}$ to $\frac{3}{4}$ inch (3 to 4 mm) long and $\frac{1}{16}$ inch (1.5 mm) in diameter, an oval oil gland being situated in the middle of the convex back of the leaf. It has a characteristic taste, and a distinctive odour when bruised, by which it can be recognised from other nearly allied species of *Juniperus*. French Savin is sometimes derived from *Juniperus phænicea*, L., and *Juniperus thurifera*, L., var. *Gallia*—Coney, *P J* (4) xvi 829-831.

Preparations

TINCTURA SABINÆ—1 of Savin Tops, dried and coarsely powdered, percolated with Alcohol (60 p c), to yield 8.

Dose—20 to 60 minims = 1.2 to 3.6 c c.

B P 1885, omitted in *B P* 1898, and now incorporated in the *B P C*.

U S has Fluidextractum Sabinæ, 1 in 1, with Alcohol (95 p c).

UNGUENTUM SABINÆ—Fresh Savin Tops, bruised, 8 Yellow Beeswax, 3, Benzoated Lard, 16, melt the Lard and the Beeswax together on a water bath, add the Savin, digest 20 minutes, strain, and press through calico.

B P 1885, omitted in *B P* 1898, and now incorporated in the *B P C*.

OLEUM SABINÆ—A colourless, or pale yellow, oily liquid, possessing a peculiar, unpleasant, narcotic odour, and bitter, pungent, camphoraceous taste. It should be preserved in dark, amber tinted, well closed bottles. It is liable to become darker in colour and to thicken on exposure to air. It is a volatile Oil distilled in Britain from fresh Savin.

The principal constituent of the Oil is an Alcohol Sabinol, which appears in the Oil chiefly in the form of Sabinol Acetate, corresponding to a content of about 40 to 44 p c. It also contains Cadinene Pinene and Camphene, and an aldehyde or ketone possessing when reformed from its Sodium Acid Sulphite compound an odour faintly resembling Cumic Aldehyde.

Solubility—4 in 1 of Alcohol (90 p c), in all proportions of Absolute Alcohol.

Dose—1 to 4 minims = 0.06 to 0.24 c c, in pill with Soap and Liquorice Powder, *see* p 897.

Foreign Pharmacopœias—Official in Belg, Jap, Port and U S.

Tests—Savin Oil has a sp gr of 0.910 to 0.930. An optical rotation of $+40^{\circ}$ to $+60^{\circ}$, a Saponification value of 115 to 125. It dissolves to form a clear solution in about half its volume or more of Alcohol (90 p c), but does not form a perfectly clear solution in from 15 to 20 volumes of Alcohol (80 p c).

SACCHARINUM. *See* GLUSIDUM

SACCHARUM LACTIS.

MILK SUGAR

B P Syn—LACTOSI

$C_{12}H_{22}O_{11}$, H_2O , eq 357.48

FR, LACTOSE, GER, MILCHZUCKER, ITAL, LATTOSIO, SPAN, LACTOSA

White, or almost white, prismatic crystals, or masses of crystals, or as a white, odourless powder, possessing a slightly sweet taste. It is obtained by recrystallisation from the evaporated Whey of cow's Milk.

Solubility.—1 in 6 of cold Water, 1 in 1 of boiling Water, almost insoluble in Alcohol (90 p c)

Medicinal Properties.—Nutrient in various cases of extreme irritability of the stomach, as it does not ferment so readily as Cane Sugar, it is used to mix with the food of children. Added to diluted cow's Milk a good substitute for human Milk is formed. Slightly diuretic in cardiac dropsy. U with potent medicinal powders, in order to equally.

Dose.—60 to 120 grains = 4 to 8 g. in more in Water

Official Preparations—Used in the preparation of *Extractum Belladonnae*, *Alcoholcum*, *Extractum Nucis Vomicae*, *Extractum Physostigmatis*, *Extractum Strophanthi* and *Pulvis Elaterini Compositus*

Foreign Pharmacopœias—Official in all. Fr (*Lactose*), Ital (*Lattosio*), Mex (*Azúcar de Leche*), Port (*Assucar de Leite*), Span (*Lactosa*)

Tests.—Milk Sugar dissolves in Water forming a clear solution, which is neutral in reaction towards Litmus paper and which is dextrogyrate. A small quantity of the aqueous solution added to Potassio-cupric Tartrate (Fehling's) Solution, produces an immediate red precipitate of Cuprous Oxide, when boiled with an equal volume of Sodium Hydroxide Solution the liquid turns yellowish-brown and finally a brownish-red. Milk Sugar may be readily determined by titration with Pavy's Copper Solution. When present together with Cane Sugar the latter may be also determined by inverting with Citric Acid and making a separate determination of the reducing power of the inverted solution by Pavy's Solution. The difference between the reducing power of the solution before and after inversion corresponds to the amount of Cane Sugar present. 49.4 parts of Cane Sugar have the same reducing power as 100 parts of Milk Sugar. Citric Acid does not invert Milk Sugar but readily inverts Cane Sugar. Milk Sugar when boiled with Sulphuric Acid undergoes hydrolysis, with the formation of Dextrose and Galactose.

The more generally occurring impurities are heavy metals, *eg*, Arsenic, Copper, Lead, Iron and Zinc, Cane Sugar, Dextrin, Starch, free Lactic Acid, and mineral matter. Arsenic, Copper, Lead, Iron and Zinc, if present, may be detected by the Hydrogen Sulphide test either in a solution rendered faintly acid with diluted Hydrochloric Acid or in one made alkaline with Ammonia Solution. Cane Sugar and Dextrin, if present, may be detected by the Alcohol test and Starch by the Iodine test. If free Lactic Acid be present, the quantity may be determined by titrating a weighed quantity of the substance with Volumetric Sodium Hydroxide Solution using Phenolphthalein Solution as an indicator of neutrality. The B.P. requires that a solution of 1 gramme in 10 cc of Water should produce a red coloration under these conditions with 3 drops of the volumetric alkali solution. Assuming that the 3 drops are required, this would be equivalent to 1.3 p c of Lactic Acid. The majority of average commercial samples require considerably less. The Alcohol and Iodine tests are described below. Mineral matter,

if present, may be detected by the residue left on ignition. The *BP* and the *USP* both require that it shall not leave more than $\frac{1}{4}$ p c of ash after ignition, the *PG* that 0.2 of a gramme of the specimen shall not leave a weighable residue.

Alcohol—If a mixture of 15 grammes of Milk Sugar and 50 c c of diluted Alcohol be allowed to stand for half an hour with occasional agitation and then filtered, a filtrate is obtained 10 c c of which mixed with an equal volume of Absolute Alcohol should not become turbid and on evaporation on a water bath should not leave more than 0.04 gramme of residue. *PG*, 1 gramme digested for half an hour, with intervals of occasional shaking, with 10 c c of Alcohol (48.9 p c), filtered, and the filtrate mixed with an equal volume of Absolute Alcohol should afford a clear mixture, which, when evaporated on a water bath, should yield not more than 0.03 of a gramme of residue, indicating the absence of Cane Sugar, *USP*.

Iodine—If 1 gramme of Milk Sugar be boiled with 50 c c of Distilled Water for 5 minutes and the solution cooled, no blue coloration should be produced upon the addition of 1 drop of T.S. of Iodine, *USP*.

SACCHARUM PURIFICATUM.

REFINED SUGAR

BP Syn—SUCROSI

$C_{12}H_{22}O_{11}$, eq 339.60

FR, SUCRE BLANC OFFICINAL, GER, ZUCKER, ITAL, ZUCCHERO,
SPAN, AZUCAR

Colourless, translucent, prismatic crystals, or a fine, white, crystalline powder, possessing a sweet characteristic taste. Permanent in the air. Obtained from the Juice of the Sugar-cane.

Solubility—100 in 45 of Water, measures 113, 1 in 100 of Alcohol (20 p c).

Medicinal Properties—Nutrient, demulcent, used in catarrhal affections in the form of Candy, Syrup, etc., also in irritant corrosive poisoning. Employed almost entirely as a sweetening agent and as a preservative, and to assist the suspension of powders. It assists the solution of Lime in Water.

It is taken as a respiratory fuel by men about to undertake excessive physical exertion.

Official Preparation—Syrupus. Sugar in some form is contained in all Syrups and Lozenges, several Confections, Mixtures, Pills and Powders.

Foreign Pharmacopœias.—Official in all except New Fr (Sucre Blanc Officiel), Ital (Zucchero), Mex (Azúcar de Caña), Port (Assucar), Span (Azucar).

Tests—Refined Sugar dissolves readily in Water, forming a clear solution which is neutral to Litmus paper, the *USP* states that the aqueous or alcoholic solution is neutral to Litmus paper, and the *PG* that aqueous and alcoholic solutions are neutral to Litmus paper. The *USP* states that an aqueous solution saturated at 25° C (77° F) should possess a sp gr of 1.340. A crystal of Sugar, when moistened with strong Sulphuric Acid, immediately chars and

swells up, forming a black mass. When a few crystals are mixed with a little powdered Potassium Chlorate and touched with a drop of concentrated Sulphuric Acid, the mixture instantly ignites.

The more generally occurring impurities are insoluble salts, ultramarine, and Prussian blue. Glucose or Inverted Sugar, Calcium, Chlorides, Sulphates, and mineral matter. Insoluble salts, ultramarine, and Prussian blue, may be detected by the Water and Alcohol test described below. Heavy metals, such as Copper, Iron, Lead, and Zinc, may be detected by the addition of Hydrogen Sulphide to either a faintly acid solution or a solution rendered alkaline by Ammonia. The official directions for the detection of Glucose or Inverted Sugar are to heat the Syrup to a temperature of about 82° C (180° F) with Potassium Cupric Tartrate (Fehling's) Solution, or with Copper Sulphate Solution and an excess of Potassium Hydroxide Solution, when not more than a trace of red or yellowish precipitate should be produced, the *USP* employs the Silver Nitrate and Ammonia test described in the small type below. The Ammonium Oxalate, Silver Nitrate, and Barium Nitrate tests described below serve to detect Calcium, Chlorides, and Sulphates if present. It should leave scarcely any ash when ignited with free access of air. The *USP* makes no reference to the amount of ash, the *PG* states 0.5 of a gramme, when ignited, should leave no weighable residue.

Water and Alcohol—One part by weight of Sugar should form, with 0.5 parts by weight of Water, a colourless, odourless syrup, possessing a purely saccharine taste and which mixes in all proportions with Alcohol (90 p.c.), *PG*.

Both the aqueous and alcoholic solutions should be clear and transparent. When kept in large, well-closed and completely filled bottles, the solutions should not deposit a sediment on prolonged standing (absence of insoluble salts, ultramarine, Prussian blue, etc.), *USP*.

Hydrogen Sulphide—An aqueous solution (1-20) should not become turbid with T.S. of Hydrogen Sulphide, *PG*.

Ammonium Oxalate—An aqueous solution (1-20) should not become turbid more than opalescent with T.S. of Ammonium Oxalate, *PG*.

Silver Nitrate—An aqueous solution (1-20) should not become more than opalescent with T.S. of Silver Nitrate, *PG*.

Silver Nitrate and Ammonia—L. . . . of Sugar be dissolved in 10 cc of boiling Water, the solution . . . 4 or 5 drops of Silver Nitrate T.S., then about 2 cc of Ammonia Water added, and the liquid quickly brought to the boiling point, not more than a slight coloration and no black precipitate should appear in the liquid after standing at rest for 5 minutes, *USP*.

Barium Nitrate—An aqueous solution (1-20) should not become more than opalescent with T.S. of Barium Nitrate, *PG*.

Preparation.

SYRUPUS. SYRUP

Dissolve 10 of Sugar in 5 of boiling Distilled Water, and finally make up the total weight to 15.

9 measures of Syrup contain 8 of Sugar.

Foreign Pharmacopœias.—Official in all, .

Tests.—Syrup has a sp gr of 1.330. It should be strongly dextrogyrate. A small quantity, when placed in a test tube with some Potassium cupric Tartrate (Fehling's) Solution, and heated in a water bath, should yield no decided red precipitate. The Syrup should be neutral in reaction towards Litmus paper.

Not Official

SALEP

The prepared tubers of *Orchis Morio*, L., and other species of *Orchis*.

Medicinal Properties—Mucilaginous and nutrient.

Foreign Pharmacopœias—Official in Austri, Belg., Dutch, Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Swed. and Swiss.

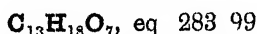
MUCILAGO SALEP—Powdered Salep 1, agitate well with cold Water 10, pour on this boiling Water 90, and stir till cold.

Foreign Pharmacopœias—Official in Dutch, Ger., Jap., Norw., Russ., Swed. and Swiss, 1 in 100.

Salib Misri, the Salep of the Indian Bazaar, is derived from a species of *Eulophia*.

SALICINUM.

SALICIN



FR., SALICINE, GER., SALICIN, ITAL., SALICINA

Colourless, tabular crystals, or slender, white, shining, acicular crystals, possessing a very bitter taste. It is a glucoside occurring naturally in the Bark and Leaves of various species of *Salix* and of *Populus*.

It should be kept in well-stoppered vessels of a dark amber tint.

Jowett (*Report of the Wellcome Chemical Research Laboratories, JCS Trans.* '00, 707, *YBP* '00, 117) states that it has been generally assumed that the different species of Willow Bark contain the same glucoside (Salicin), but having had occasion to examine a bark purchased as Black Willow he found that the crystalline principle obtained by the usual method for preparing Salicin was not that substance, but a new glucoside for which the name of Salinigrin was proposed. During the determination of the constitution of the new glucoside he established the interesting fact that, whilst Salicin is the glucoside of *o*-Hydroxybenzyl Alcohol, Salinigrin is the glucoside of *m*-Hydroxybenzaldehyde. Salinigrin can easily be distinguished from Salicin by affording a colourless solution with Sulphuric Acid, whilst Salicin, under similar conditions, produces a blood red colour. The investigations by Jowett into the variations in the occurrence of Salicin and Salinigrin (*CD* '02, 11347, *YBP* '02, 490) in Willow and Poplar Barks, wherein a considerable number of authentic specimens

of *Salix* and *Populus* from the chief European and American species were examined, showed that, of the 33 samples examined, Salicin was only found in 1, *Salix discolor*, Muhl., that the amount of Salicin contained in the bark of the Willow or Poplar depends not only on the species, but on the season of the year at which it is collected, the sex of the tree, and possibly other factors, that the results of the investigation have shown that for practical purposes chemical assay alone can decide whether a Willow bark does or does not contain Salicin

Solubility.—1 in 28 of Water, 1 in 82 of Alcohol (90 p c), insoluble in Ether

Medicinal Properties.—Antipyretic, antiperiodic, tonic, and bitter stomachic, has been specially recommended in acute rheumatism. For the latter purpose it has been largely replaced by Sodium Salicylate, the action of which is more powerful, though not so well sustained, as Salicin, but the Salicylate has a greater tendency to cause cardiac depression, and is not so well tolerated by the stomach as Salicin. Has been recommended for the prevention and cure of influenza

20-grain doses 3 times a day given with great success in a case of lupus erythematosus —*L* '02, ii 157

15-grain doses in psoriasis the patches became paler, the scales more detachable and soon ceased to reform, while patch cleared in the centre, and finally the circle broke up —*B M J* '03, i 656, *L* '95, i 1421, '03, i 784

15 grains every 4 hours relieved the irritation and arrested maturation of the vesicles in smallpox —*B M J* '00, i 16, 512, 1387, '00, ii 127, '02, ii 179, *P. J* '02, ii 113

Dose.—5 to 20 grains = 0.31 to 1.3 grammes

Prescribing Notes.—It is given in cachets. A good pill can be made by adding 'Diluted Glucose,' q s

Effervescent Granules can be obtained containing 5 grains in each drgm

Not Official—Saligenin, Salnigrin and *Salix Nigra*

Foreign Pharmacopœias—Official in Ital, Mex, Port and U S

Tests.—Salicin melts, when pure, at 200° C (392° F). The U S P gives the m p as 201.4° C (394.5° F), when still more strongly heated [240° C (464° F)] it decomposes. When moistened with Sulphuric Acid it is coloured red. When heated with a small quantity of Potassium Bichromate and a little Sulphuric Acid it evolves an odour of Salicylic Aldehyde, recalling the odour of Meadow-Sweet. When warmed in a test-tube until it turns brown the residue, when mixed with Water, yields, on the addition of Ferric Chloride T S, a violet coloration. When evaporated to dryness with a few drops of Nitric Acid, and the yellow residue is treated with Ammonia Water, it yields, when heated on a water-bath with a small quantity of Potassium Cyanide, a blood-red coloration. It dissolves in Water, forming a solution which is colourless and neutral in reaction towards Litmus, and is not decomposed by levogyrate. The U S P states that, when moistened with Sulphuric Acid containing a trace of Molybdic Acid, a violet coloration changing to deep brownish-red is produced,

and that Sulphuric Acid containing a trace of Potassium Iodide Solution yields a dark red coloration, changing to deep purple, and that Sulphuric Acid containing about one-fifth of its volume of Formaldehyde Solution produces a deep purplish-red colour. Salicin may be distinguished from alkaloids by yielding, when dissolved in Water, neither a precipitate with Potassium mercuric Iodide (Mayer's) Solution, nor with Tannic nor with Picric Acid Solutions, nor the other usual reagents for alkaloids. When ignited with free access of air it should leave no residue, the latter requirement is common to both the *BP* and the *USP*.

Not Official

SALIGENIN—Small, tabular crystals, having a very faint, sweetish taste, soluble in Water, readily soluble in Alcohol (90 p c) and in Ether. It is obtained by the action of Formic Aldehyde on Phenol in alkaline solution, or by the action of emulsin or of diluted mineral acids on Salicin. It has been recommended in acute rheumatism and in gout.—*P J* (3) xxv 755, 1115, '95, ii 175. **Dose**—4 grains = 0.26 grammes.

SALINIGRIN—Was found by Jowett (*J C S Trans* '00, 707) in an examination of a bark purchased as that of Black Willow, but which could not be, however, identified then as other than some species of *Salix*, the crystalline substance possessed a m p of 193° C (379.4° F) Cor. Its aqueous solution had an optical rotation of -85°, the substance gave no coloration with Sulphuric Acid, on hydrolysis it yielded a crystalline Meta hydroxybenzaldehyde.

SALIX NIGRA—The Bark has been recommended as a sexual and general sedative.—*B M J* '87, ii 237, *L* '88, i 369.

The dose of the Fluid Extract (1 in 1) is 30 to 60 minims = 1.8 to 3.6 cc.

SALOL.

PHENYL SALICYLATE

$C_7H_5O_3$, C_6H_5 , eq 212.47

Fr, SALICYLATE DE PHENYLE, GER, PHENYLSALICYLAT, ITAL, SALOLO, SPAN, SALICILATO DE FENOL.

Colourless, translucent, needle-shaped crystals, or a white crystalline powder, possessing a peculiar and characteristic aromatic odour, and but a slight taste. It is the Salicylic Ester of Phenyl.

Solubility—1 in 12 of Alcohol (90 p c), 2 in 1 of Ether, 3 in 1 of Chloroform, 1 in 4 of Almond Oil, 1 in 10 of Liquid Paraffin. Insoluble in cold Water.

Medicinal Properties—Antipyretic, antiseptic, and intestinal disinfectant. It passes through the stomach unchanged, and is decomposed into Carbolic and Salicylic acids by the alkalinity of the small intestine. It has been recommended in acute and chronic rheumatism, in cholera, in typhoid fever, in intestinal tuberculosis and in smallpox. One of the best antiseptics for intestinal dyspepsia and fermentation. Useful also as a urinary antiseptic. When given in excessive doses, or repeated frequently, has

given rise to toxic symptoms. Externally it is used for the same purposes as Iodoform

Combined with a blood tonic in anæmia, *MA* '95, 103, and pernicious anæmia, *L* '94, ii 1274, in diarrhoea of phthisis, *Pr* lin 275, in choleraic diarrhoea, *TG* '94, 40, good result in gonorrhoea, *L* '90, i 644, an intestinal and urinary disinfectant *BMJ* '93, i 643

Owing to its low m.p. (about $108^{\circ}\text{F} = 42^{\circ}\text{C}$) it is useful in filling up irregular or superficial bone cavities, also as a stopping for carious teeth — *BMJE* '96, i 64, *PJ* '95, ii 216

Formation of Salol calculus from its internal administration — *BMJ* '97, ii 78, *PJ* '97, ii 446

Has been used (*L* '04, ii 1209) in 10-grain doses as an internal antiseptic in ulcerative colitis

10 to 20-grain doses, either alone or in emulsion, with Castor Oil and Gum Acacia as an intestinal antiseptic in dysentery — *IMG* '05, ii 281

In compressed form it is less reliable than when given in suspension in Petroleum Emulsion — *BMJ* '05, ii 1703

Dose.—5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes—*It is given in cachets, mixtures, powders, or compressed tablets. In mixtures it should be suspended with Compound Tragacanth Powder, but it is best dissolved in a fixed Oil, and emulsified by Gum Acacia (see below Emulsion Salol). Salol with $\frac{1}{2}$ of Compound Tragacanth Powder, will make a good pill with 'Diluted Glucose'*

A good mouth-wash can be made by dissolving 60 grains of Salol in 6 fl oz of Alcohol (90 p.c.), and adding 10 minims of Oil of Peppermint and 5 minims of Oil of Anise. It can, if desired, be sweetened with an addition of $\frac{1}{16}$ grain of Saccharin

Not Official—Emulsion Salol, Pommade de Salicylate de Phényle, Salol Camphor Salol Mouth-wash, Salol Varnish for Pills, and Salophen

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fl, Ger, Ital, Jap, Mex, Norw, Russ, Span, Swed, Swiss and U.S.

Tests.—Commercial Salol melts at 41.37°C (106.46°F), dried Salol melts at 42.53°C (108.55°F), purified Salol melts at 42.47°C (108.44°F). The *BP* gives the m.p. as 42° to 43°C (107.6° to 109.4°F). The *USP* and the *PG* 42°C (107.6°F). It dissolves readily in Alcohol, the alcoholic solution being neutral in reaction towards Litmus paper, and yielding on the addition of Bromine Solution a white precipitate, and on the addition of Ferric Chloride T.S. a violet coloration. The *USP* and the *PG* both employ diluted Ferric Chloride Solution in carrying out this test. When dissolved in a little warm Sodium Hydroxide Solution it yields when cooled and acidified with diluted Sulphuric Acid a white precipitate, and on warming, an odour of Phenol. The *BP* states that when Salol is melted with Sodium Hydroxide and acidulated with Hydrochloric Acid a white precipitate is produced and Phenol is evolved. The object in using solid Sodium Hydroxide, when Potassium or Sodium Hydroxide Solution is equally convenient and far less troublesome, is not apparent. The separated Salicylic Acid, when washed and carefully dried should possess the m.p. and answer the tests distinctive of Salicylic Acid given under Acidum Salicylicum.

Bearing in mind the fact that Phenol forms no chemical combination with Sodium Hydroxide in the true sense of the term, any alkali Hydroxide being titratable with Normal Acid Solution as if no

Phenol were present, and with a standard acid forms a salt neutral to Phenolphthalein Solution, the author has suggested (*P J* '05, 1720) a convenient method for approximately determining the purity of a salt by the Saponification equivalent -- A weighed quantity of 0.5 of a gramme of the sample is carefully weighed out into a flask, 5 c.c. of Normal Volumetric Sodium Hydroxide Solution added, and 10 c.c. of Water, the mixture saponified on a water bath, it is cooled, the excess of Normal Volumetric Sodium Hydroxide Solution titrated with Tenth normal Volumetric Sulphuric Acid Solution. A carefully measured 5 c.c. of the Normal Alkali Solution is treated in an exactly similar manner as a blank experiment, the difference of the number of c.c. of Tenth normal Volumetric Sulphuric Acid Solution required for the specimen and that required in the blank experiment is calculated into terms of Phenyl Salicylate 1 c.c. of Tenth normal Sodium Hydroxide Solution absorbed represents 0.005571 gramme of Potassium Hydroxide or 0.021247 gramme of Phenyl Salicylate.

The more generally occurring impurities are uncombined Salicylic Acid or Phenol, Sulphates and Chlorides. The free acid, if present, may be detected by the behaviour of the sample or its solution towards blue Litmus paper. When shaken with 50 times its weight of Water and filtered the filtrate should neither afford a blue nor a violet coloration with diluted Ferric Chloride T.S. (the *B.P.* says Ferric Chloride Solution), nor should it produce a turbidity with Silver Nitrate or Barium Chloride Solutions. When heated with free access of air it should leave no weighable residue. The *P.G.* states that 0.1 of a gramme when ignited shall leave no weighable residue.

Ferric Chloride -- If 1 gramme of Phenyl Salicylate be shaken with 50 c.c. of Water and filtered, and if 5 drops of Ferric Chloride T.S., previously diluted with 20 volumes of Water be added to the filtrate, the latter should show either no colour or at most a trace, *U.S.P.* Also given in the *P.G.*, which states that the filtrate should not be affected.

Barium Nitrate -- A filtrate obtained as above should be unaffected by T.S. of Barium Nitrate, *P.G.*, should show no turbidity, *U.S.P.*

Silver Nitrate -- Another portion of a filtrate as above should be unaffected by T.S. of Silver Nitrate, *P.G.*, should show no turbidity, *U.S.P.*

Not Official

EMULSIO SALOL -- Salol, 40 grains, Almond Oil, 4 fl. drms., Powdered Gum Acacia, 120 grains, Syrup, 2 fl. drms., Peppermint Water, to 2 fl. oz.

POMMADE DE SALICYLATE DE PHÉNYLE -- Phenyl Salicylate, 1, Vaseline, 9 -- *F.*

SALOL CAMPHOR -- Prepared by moistening 1 of Camphor with Alcohol and titrating it with $1\frac{1}{2}$ Salol till a transparent liquid is obtained. Has been found useful in treatment of furuncles and carbuncles -- *B.M.F.E.* 95, 1181.

SALOL MOUTH-WASH -- Salol, 60 grains, Oil of Peppermint, 10 minims, Oil of Anise, 5 minims, Alcohol (90 p.c.), to 6 fl. oz. It can, if desired, be sweetened with an addition of $\frac{1}{16}$ gram of Saccharin.

A modification appears in the *B.P.C.*, under the title **Liquor Salolis Compositus**, with the synonym Salol Mouth Wash as follows --

Salol, 2.50, Thymol, 0.25, Spirit of Anise, 1, Oil of Peppermint, 0.50, Elixir of Gluside, 2.50, Alcohol, q.s. to produce 100

SALOL VARNISH FOR PILLS—Salol, 2, Shellac, 3, Absolute Alcohol, 3, Ether, 3—*Martindale*

This has been incorporated in the *BPC* under the title **Solutio Salolis Etherea**, with the synonym Salol Pill Varnish as follows—

Salol, 20, Shellac, 80, Ether, 30, Absolute Alcohol, *q s* to produce 100

SALOPHEN Acetylparamidophenol Salicylate $C_{14}H_{11}NO_3$, eq 269.11—
A white, crystalline powder, insoluble in Water, soluble in Alcohol and in Ether.
It appears in the *Fr Codex* (1908) under the title of Acétylpara-amino-salol

Medicinal Properties—Analgesic and antipyretic. Has been recommended in acute and subacute rheumatism, and in neuralgia

Dose—10 to 30 grains = 0.65 to 2 grammes, usually given in cachets

Official in Belg, Fr, Mex, Swed and Swiss

Tests—Salophen melts at 187° to 188° C (368.6° to 370.4° F), *Fr Codex* gives 188° C (370.4° F). It dissolves in Alcohol, forming a solution which is faintly acid towards Litmus paper, and which is coloured violet by the addition of Ferric Chloride T.S., and which produces a voluminous white precipitate with Bromine Solution. It dissolves without change of colour in concentrated Sulphuric Acid. Potassium or Sodium Hydroxide Solution readily decomposes it into Salicylic Acid and Acetyl-para-amidophenol. The *Fr Codex* includes a test for the acetyl radicle, requiring that when moderately heated with a mixture of Alcohol and Sulphuric Acid it shall evolve an odour of Acetic Ether. The liberated Salicylic Acid, when separated, washed and carefully dried, should possess the m.p. and respond to the tests characteristic of Salicylic Acid given under Acidum Salicylicum

SAMBUCI FLORES.

ELDER FLOWERS

Fr, SUREAU, GER, HOLUNDERBLUTHEN, ITAL, SAMBUCCO, SPAN, SAUCO.

The Flowers of *Sambucus nigra*, L, separated from the stalks

Descriptive Notes.—The official description does not specify whether the Elder flowers should be fresh or not, but under *Aqua Floræ Sambuci* the fresh flowers are ordered. The flowers should be separated from the flower stalks. In the West of England the dried flowers are commonly sold as a remedy for catarrh, the whole inflorescence being dried. The flowers readily blacken if left in heaps, and need to be quickly dried in a current of warm air in order to keep their colour. The small rotate corolla is nearly white when fresh, but dull yellowish-white when dried. The anthers are yellow, in the only other British species, *S. Ebulus*, they are pink. The fresh flowers have a slightly bitter taste and a faint but characteristic odour. In the United States the allied species *S. Canadensis*, L., was official, and is still in use. It differs from the British species chiefly in the leaves having 3 to 4 pairs of leaflets and in being sometimes bipinnate, but the flowers present no marked difference.

Official Preparation—*Aqua Sambuci*

Not Official—*Unguentum Sambuci*, *Unguentum Sambuci (Viride)*

Foreign Pharmacopœias.—Official in all except U.S. Fr (Sureau) Ital (Sambucco), Mex and Span (Saucó), Port (Sabugueiro)

Preparation**AQUA SAMBUCI** ELDER FLOWER WATER

Fresh Elder Flowers, 1 (or an equivalent quantity of the Flowers preserved whilst fresh with Common Salt), Water, 5, distil 1
(1 in 1)

Chiefly used for lotions and collyria

Not Official

UNGUENTUM SAMBUCI —Elder Flowers, fresh, Lard, of each, 16 or Boil the Elder Flowers in the Lard until they become crisp, then press through a linen cloth —*P L* 1851

This has been incorporated in the *B P C*

UNGUENTUM SAMBUCI (VIRIDE) —Elder Leaves, fresh, 3, Prepared Lard, 4, Prepared Suet, 2, boil the leaves with the Lard till they become crisp, strain, express, add the Suet and melt them together —*Dublin Pharm*

This has been incorporated in the *B P C*

Foreign Pharmacopœias —Official in Port 1 in 4

SANTALI OLEUM.

OIL OF SANDAL WOOD

B P Syn —OIL OF SANDAL WOOD

FR, ESSENCE DE SANTAL, GFR, SANDLÖL ITAL, ESSENZA DI SANDALO, SPAN, ESSENCIA DE SANDATO

A pale yellow, or yellow, somewhat viscid, oily liquid, having a characteristic, persistent, aromatic odour, and unpleasant, nauseous taste

It should be kept in well-closed bottles of a dark amber tint in a cool atmosphere and protected as far as possible from contact with the light

It is distilled from the Wood of *Santalum album*, L

East Indian Sandal Wood Oil is alone official in the *B P*, the *USP* requires the Oil to contain not less than 90 p c of alcohols calculated as Santalol, the *P G* does not state the necessary proportion of Santalol

The chief constituent of the Oil is an alcohol Santalol, which is capable of determination by acetylation

Solubility.—In less than its own weight of Alcohol (90 p c)

Medicinal Properties —A stimulating disinfectant to the mucous membranes of the bladder and urethra, and also of the bronchial mucous membrane, prescribed extensively for subacute and chronic gonorrhœa, it is best taken about an hour and a half after meals

Dose.—5 to 30 minims = 0.3 to 1.8 cc

Prescribing Notes —Generally given in capsules or in a mixture suspended with Mucilage of Acacia, or Tragacanth It is best taken in Capsules, as the taste is nauseous Sometimes prescribed with Buchu and Cubebs

Not Official —Capsules of Sandal Oil, Liquor Santali Compositus, Mistura Olei Santali, Mistura Santali Composita, Mistura Santali Composita cum Morphina, Mistura Olei Santali, Gonal and Santyl

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Ger, Jap, Norw, Span, Swiss and U S

Tests—Sandal Wood Oil has a sp gr of 0.975 to 0.985. The *B P* states 0.975 to 0.980, the *U S P* 0.965 to 0.980 at 25° C (77° F), the *P G* 0.975 to 0.985. It is officially required to dissolve in six times its volume of Alcohol (70 p c), to form a clear solution, but it must be remembered that the solubility of the Oil decreases with age and that an Oil which has been kept a considerable time or which has been badly preserved may not give a clear solution. The *U S P* requires that it should dissolve in 5 volumes of Alcohol (70 p c) presumably at 25° C (77° F). The *P G* is more specific with regard to the temperature at which solution in this volume of Alcohol is required to take place, and states that it shall dissolve in 5 parts, by weight, of Alcohol (68 to 69 p c) at 20° C (68° F) to form a clear solution possessing a faintly acid reaction. The temperature at which solution is required to be effected should have been mentioned in the *B P*, as it makes a considerable difference whether the solubility figure is determined at 15.5° C (60° F), 20° C (68° F) or 25° C (77° F). It is levogyrate, the optical rotation being from -16° to -20° in a tube of 100 mm length. These are the figures required by the *B P*. The *U S P* requires that the optical rotation should not be less than -16° nor more than -20° in a 100 mm tube at a temperature of 25° C (77° F). The *P G* does not state the optical rotation. Considerable controversy has raged round the question of the optical rotation, it being contended that specimens of undoubted purity (English distilled) are occasionally outside these limits, and rotations of -14° to -22° have been recorded. The majority of evidence appears to be in favour of the -16° to -20° limit, it not being considered a good policy to widen the official limits in order to include a few exceptional oils possessing a rotation outside the above limits. The optical rotation has been considered fallacious in judging the purity of an Oil, but it must be recollected that it frequently supplies important information as to the nature of the substance with which an Oil is adulterated. The refractive index should be not below 1.503. The alcohol-content of the Oil calculated in terms of Santalol should not fall below 90 p c. Neither the *B P* nor the *P G* gives a requisite Santalol content, the *U S P* requires that it shall contain not less than 90 p c of alcohols calculated as Santalol, as determined by the process given in small type below. A Santalol determination figure, possibly 94 p c as a minimum, has been suggested for inclusion in the *B P*. All authorities are agreed that the Santalol content for a genuine Oil should not fall below 90 p c, but the majority consider that although the total amount of Santalol present in Oils of undoubted purity is below 94 p c, the standard is somewhat too high for practical purposes and that it would be wiser to adopt the standard recommended by Parry and Schimmel of at least 90 p c. Besides the alcohol, Santalol, the Oil also contains esters of that Alcohol present chiefly in the form of Acetate, their percentage varies from 2 to 6 p c and they may be determined by saponifying a known weight of the Oil with Semi-

normal Volumetric Alcoholic Potassium Hydroxide Solution, the excess of the latter solution with Semi-normal Sulphuric Acid Solution, using Phenolphthalein Solution as indicator of neutrality. The number of c.c. of Semi-normal Volumetric Alcoholic Potassium Solution absorbed by the Oil multiplied first by 0.1301 and then by 100, and the product divided by the weight of Oil taken, yields the percentage of esters in terms of Santalol Acetate. Neither the *B.P.*, the *U.S.P.* nor the *P.G.* includes figures for an ester content. In carrying out the volumetric determination of Santalol content, in the place of washing the acetylated Oil with Water, whereby, owing to the formation of an emulsion, a fair quantity of the Oil is lost, it has been suggested (*Proc. Amer. Pharm. Assoc.* '06, 887) that a 10 p.c. Sodium Chloride Solution should be used. The formula by which the Santalol content should be calculated is also a matter of importance. Schimmel states that Santalol is correctly represented by the formula $C_{15}H_{24}O$, which is the only one which ought to come under consideration.

The more generally occurring impurities are Oils derived from other varieties of Sandal Wood, Cedar Wood Oil, Castor Oil, or other fixed Oils and Rosin. The *U.S.P.* includes a test for chlorinated products, which is described under Silver Nitrate. The solubility of the Oil in Alcohol (70 p.c.) detects the presence of Castor Oil or other fixed Oils or West Indian Sandal Wood Oil. Cedar Wood Oil, Castor Oil or fixed Oils and Rosin may also be detected by the Acid and Ester values and the decrease in the optical rotation, as well as by the diminution in the percentage of Santalol. It is stated (*Analyst*, '95, 174) that genuine Sandal Wood Oil gives with Bromide of Tin (*see* Oleum Lini) a red coloration, whilst West Indian Sandal Wood Oil gives a blue or a green colour. The paucity of information contained in the official monograph suggests a recommendation to the effect that the *B.P.* monograph requires revision.

Silver Nitrate - If a small strip of filter paper folded in the form of a taper and saturated with Oil of Santal be placed in a small porcelain dish, and a clean beaker moistened on the inner surface with Distilled Water be inverted over the small dish immediately after igniting the taper, a part of the products of combustion will be absorbed by the Water, if the beaker be then rinsed with a little Distilled Water and the liquid filtered, the filtrate should yield no turbidity upon the addition of a few drops of Silver Nitrate T.S., *U.S.P.*

Volumetric Determination of Santalol - A measured quantity of 10 c.c. is introduced into an acetylation flask, together with 10 c.c. of Acetic Acid Anhydride, and about 2 grammes of anhydrous Sodium Acetate, and the mixture is boiled gently for 1½ hours, when cool the acetylated Oil is washed first with Distilled Water and subsequently with Sodium Hydroxide T.S., until the mixture is faintly alkaline to Phenolphthalein T.S., and it is then dried by means of fused Calcium Chloride. Filter and transfer 3 c.c. of the dried acetylated Oil into a flask having a capacity of 100 c.c., and after having ascertained accurately the weight, saponify by boiling gently for 1 hour under a reflux condenser with 50 c.c. of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution, titrating the excess of the latter with Semi-normal Volumetric Sulphuric Acid Solution, employing Phenolphthalein Solution as an indicator of neutrality. The number of c.c. of Semi-normal Volumetric Sulphuric Acid Solution required is subtracted from 50, the difference is multiplied by 11.026 and the product divided by the weight of the dry acetylated Oil employed

(minus the number of cc of Semi-normal Volumetric Alcoholic Potassium Hydroxide Solution absorbed by the acetylated Oil multiplied by 0.021), the quotient corresponds to the p.c. of Santalol present in the sample—*U S P*

Not Official

CAPSULES OF SANDAL OIL—Containing 10 and 20 minims in each

The Oil used in these capsules is frequently adulterated, Castor Oil, flavoured with Sandal Wood Oil, has been used for this purpose, but, of course, is readily detected. The favourite adulteration is Oil of West Indian Sandal Wood, this has been reported (*C D '06*, 1 211) in specimens of capsules manufactured in London, it may be recognised by tests given under *Oleum Santali*

LIQUOR SANTALI COMPOSITUS—Sandal Wood Oil, 1, Soluble Solution of Copaiba, 22, Spirit of Cinnamon, $\frac{1}{2}$, Tincture of Buchu (1 in 5 S V R), $8\frac{1}{2}$, Tincture of Cubebs (1 in 5 S V R), 3—*Pharm Form*

Oil of Sandal Wood, 5, Spirit of Cinnamon, 2 50, Tincture of Buchu, 17, Tincture of Cubebs, 15, Alcohol, *q s* to produce 100—*B P C*

MISTURA OLEI SANTALI—*Oleum Santali*, 30 minims, Mucilage of Acacia, 1 fl drm, Syrup, 1 fl drm, Tincture of Orange, 30 minims, Water, to 1 fl oz, for a dose 3 times a day—*Squre*

Mistura Olei Santali—Oil of Sandal Wood, 15 minims, Mucilage of Gum Acacia, 30 minims, Cinnamon Water, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

MISTURA SANTALI COMPOSITA—Sandal Wood Oil, $12\frac{1}{2}$ drm, Oil of Cassia, $1\frac{1}{2}$ drm, Oil of Pimento, 40 minims, Rectified Spirit, $8\frac{1}{2}$ oz (Nisbet's Specific)—*Pharm Form*

Oil of Sandal Wood, 30, Oil of Cassia, 3 50, Oil of Pimento, 1 50, Alcohol, *q s* to produce 100—*B P C*

MISTURA SANTALI COMPOSITA CUM MORPHINA—Sandal Wood Oil, 4 oz, Oil of Pimento, 4 drm, Oil of Cassia, 2 drm, Morphine, 9 grains, Rectified Spirit, to produce 12 oz (Nisbet's Specific)—*J. J. I. R. M.*

Oil of Sandal Wood, 85, Oil of Cassia, 2 25, Oil of Pimento, 4 50, Morphine Hydrochloride, 0 15, Alcohol, *q s* to produce 100—*B P C*

GONAL—A colourless, oily liquid, sp gr 0.978 to 0.980, containing the alcohol constituents of Sandal Wood Oil. It has a faint odour of the latter. An irritating sesquiterpene Santalene is stated to be removed during its preparation. Introduced as a purified Sandal Wood Oil, and recommended for urethritis and gonorrhoea. **Gonoral** was a somewhat similar preparation—*B M J '01*, 1. 1407, '01, 11 512, *P J '99*, 11 34, '00, 1 333

SANTYL—A clear, yellow fluid of an oily consistency, possessing a faint odour and taste of Sandal Wood. It is insoluble in Water, but dissolves in Alcohol (90 p.c.) and in Ether. It is to be a neutral Santalol Salicylic Ester, and to contain 60 p.c. of Santalol. It was introduced as a urinary antiseptic, and is stated to be of value in acute gonorrhoea and its complications, being comparatively free from the somewhat objectionable odour and taste of Sandal Wood Oil. It is stated not to cause eructations, nor to impart a Sandal Wood odour to the breath. It may be given in doses of 30 drops taken 3 times daily, preferably in Milk or in the form of Capsules, 2 capsules being taken 4 times a day.

SANTONINUM.

SANTONIN

$C_{15}H_{18}O_3$, eq 244.29

FR, SANTONINE, GER, SANTONIN, ITAL, SANTONINA, SPAN, SANTONINA.

Colourless, odourless, pearly, hexagonal prisms, possessing a faint bitter taste. It is a crystalline principle, which is prepared from *Santonica*, or Worm-Seed, the dried, unexpanded Capitula of

Flower-Heads of *Artemisia maritima* The *USP* describes it as an inner Anhydride or Lactone of Santonic Acid obtained from Santonica

It should be kept in well closed bottles of a dark amber tint and protected as far as possible from the light, as it acquires, when exposed to the light, more particularly to direct sunlight, a yellow colour

Solubility—Sparsingly in Water, 1 in 350 of boiling Water, 1 in 50 of Alcohol (90 p c), 1 in 4 of boiling Alcohol (90 p c), 1 in 160 of Ether, 1 in 2 of Chloroform, about 1 in 400 of Olive Oil, slightly in Glycerin and in Solution of Potassium Hydroxide

Medicinal Properties—Anthelmintic Useful both for round worms and thread-worms It frequently affects the vision, causing all objects to appear yellow or green, to avoid thus unpleasantness, Santonin is given at night, the disturbance of vision then remains only for half an hour or so, after the patient awakes in the morning

Apart from its tenicide action, it is stated to possess valuable antispasmodic properties—*P J* '04, ii 967, *C D* '04, ii 1052 It is useful in certain nervous affections, in epilepsy, and against tabetic pains

A case of a child age $3\frac{1}{2}$ years, is recorded, in which, after it had received at intervals during 40 hours several 'worm lozenges' containing Santonin, a fatal issue ensued A little over 1 grain of Santonin had thus been taken, or about half the maximum dose for a child of 2 years per day Other cases have been recorded in which equally small doses have produced toxic effects—*Edin Med Jour* '08, i 188

Dose—2 to 5 grains = 0.13 to 0.32 gramme

Ph Ger maximum single dose, 0.1 gramme, maximum daily dose, 0.3 gramme

Prescribing Notes—About 3 doses are sufficient, 1 every other night followed by a brisk cathartic the morning after each dose The suppository is useful in thread-worms

Castor Oil has been recommended as a solvent for Santonin, but it will not dissolve 1 in 100, even if heat be applied, part of the Santonin will crystallise out on cooling

Official Preparation—Trochiscus Santonini

Not Official—Suppositorium Santonini, Pulvis Santonini Compositus In fantilis, Pulvis Santonini et Scammonii, and Artemisin

On account of the similarity in crystalline form, and in consequence of several accidents due to the contamination of Santonin with Strychnine, *Ger* and *US* include a test for the latter substance

Foreign Pharmacopœias—Official in all

Tests.—Santonin melts at 170° C (338° F), and if cautiously heated it may be sublimed unchanged, the *USP* gives the m p as 170.3° C (338.5° F), the *PG* 170° C (338° F) When more strongly heated it acquires a reddish-brown colour, evolving white fumes It is soluble in Potassium or Sodium Hydroxide Solutions, and when added to a warm alcoholic Solution of the former it yields a violet-red coloration It dissolves in Alcohol, the solution being lævogyrate and neutral in reaction towards Litmus paper When moistened with Sulphuric Acid or Nitric Acid no coloration is produced It is insoluble in diluted mineral acids If 0.1 of a gramme

be shaken with 1 c c of a cold mixture of Sulphuric Acid, and 1 c c of Water, no coloration should be produced, but on heating to 100° C (212° F), the addition of a drop of diluted Ferric Chloride T S a purple-violet coloration is produced, changing to brown on long continued heating. Crystals of Santonin are somewhat similar in appearance to Strychnine, and in fact have been mistaken for that substance, and tests for Strychnine, Brucine and alkaloids have been inserted in the *USP* and *PG*. The test for Strychnine and Brucine is described under the heading of Potassium Bichromate, and the test for other alkaloids under the heading of Mercuric Potassium Iodide or Iodine in small type below. When ignited with free access of air it should leave no residue, indicating the absence of mineral matter. The *USP* states that when ignited it is consumed, leaving no residue, and the *PG* that 0.2 of a gramme of the substance shall leave no residue when ignited.

Potassium Bichromate—If Santonin be boiled with 70 c c of Water and 5 parts diluted Sulphuric Acid, after cooling and filtering it gives a filtrate without any bitter taste, and in which the addition of a few drops of Potassium Bichromate T S does not produce a precipitate, *PG*.

Mercuric Potassium Iodide or Iodine—If 2 grammes of Santonin be boiled with 80 c c of Water and 5 c c of Diluted Iodine, and the liquid, after frequent shaking be allowed to become filtered, Mercuric Potassium Iodide T S or Iodine T S, should produce no cloudiness in 10 c c or the filtrate, mixed with 10 c c of Distilled Water, even after standing for 3 hours (absence of alkaloids), *USP*.

Preparation.

TROCHISCUS SANTONINI. SANTONIN LOZENGE.

1 grain of Santonin in each lozenge, with Simple Basis

Dose.—1 to 5 lozenges

Foreign Pharmacopœias—Official in Austri, Belg, Dutch, Ger, Ital., Mex, Norw, and Swiss, each containing $\frac{3}{4}$ grain, Dan, Russ, Swed. and U S $\frac{1}{2}$ grain, Fr and Port, $\frac{1}{3}$ grain, Jap and Span, $\frac{1}{3}$ grain. Not in Hung.

Not Official

SUPPOSITORIUM SANTONINI—Santonin 3 grains, with Oil of Theobroma

PULVIS SANTONINI COMPOSITUS INFANTILIS—Santonin, 1 grain, Calomel, $\frac{1}{2}$ grain, Compound Powder of Scammony, $2\frac{1}{2}$ grains—*London*

PULVIS SANTONINI ET SCAMMONII.—Santonin, 1 grain, Compound Powder of Scammony, 2 grains—*Victoria*

Artemisin (Oxysantonin) occurs in colourless crystals, becoming yellow on exposure to light, and is extracted from the mother liquor after separating the Santonin—*P J '02*, 1 294, 489, *C D '02*, 1 14

15 grains given in 3 doses at intervals of 3 hours to relieve the lightning pains of tabes—*B M J E '01*, 1 80, *T G '01*, 613

SAPO ANIMALIS**CURD SOAP**

FR, SAVON ANIMAL GER, HAUSEHLE, LIAT, SEIFEN ANIMALI,
SPAN, JABON ANIMAL

A white or whitish solid, possessing a characteristic appearance, dry and saponaceous to the touch. It is prepared by the saponification of a purified animal fat with Sodium Hydroxide.

Sapo Animalis is described in the *P U* under the title of Sapo Medicatus, it is not official in the *U S P*.

For the purpose of powdering it is not affected injuriously by drying at a temperature of 212° F (100 C).

Solubility—Sparingly in Water, 1 in 1½ of boiling Water, partially in Alcohol (90 p c), almost entirely, 1 in 2 of boiling Alcohol (90 p c).

Official Preparations—Used in the preparation of Extractum Colocynthis Compositum, Linimentum Potassii Iodidi cum Sapone, and Pilula Scammonii Composita.

Foreign Pharmacopœias—Official in Austr (Sapo Medicinalis), Belg (Sapo Stearicus), Norw (Sapo Butyraceus), Fr, (Savon Animal), Hung (Sapo Albissimus Diognistarium) Ital (Sapone Animale), Port (Sabo Animal), Russ (Sapo Sebacinus), Mex and Span (Jabon Animal) Swiss (Sapo Stearinicus), Ger, Jap and Russ (Sapo Medicatus), made with Lard and Olive Oil.

Emplastrum Saponis Formerly made with Curd Soap, now made with Hard Soap. See Sapo Durus.

Tests—Curd Soap dissolves sparingly in Water and readily in boiling Water, is sparingly soluble in Alcohol (90 p c). The *B P* requires that it shall contain no free alkali Hydroxide as determined by dissolving a weighed quantity of 5 grammes of the dried and powdered Soap in boiling Alcohol (90 p c), filtering whilst hot and washing the filter with boiling Alcohol (90 p c), using Phenolphthalein Solution as an indicator of neutrality. It is officially required to contain not more than 0.3 p c of alkali (Sodium) Carbonate as determined by dissolving in Water the residue resulting from the filtration of the boiling alcoholic solution of the Soap and washing with boiling Alcohol (90 p c). This aqueous solution is titrated with Tenth-normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality, not more than 3 c c of the Tenth-normal Sulphuric Acid Solution shall be required, 1 c c of the latter solution is equivalent to 0.00526 gramme of anhydrous Sodium Carbonate. The use of Phenolphthalein Solution as an indicator of neutrality will necessitate the boiling of the solution to dispel the Carbonic Anhydride. It would have been preferable to have used Methyl Orange Solution as an indicator, when the titration could have been carried out directly. If it is desired to ascertain the amount of alkali combined with the fatty acids in the form of a Soap, a few drops of Lacmoid Solution may be added and the titration continued until a red coloration is produced. The number of c c. of Tenth-normal

Volumetric Acid Solution used may be calculated into Sodium Oxide, 1 c.c. of Tenth-normal Volumetric Sulphuric Acid Solution is equivalent to 0.00308 gramme of Sodium Oxide. The *P.G.* requires that a solution of 1 gramme of Soap and 5 c.c. of Alcohol (90 p.c.) shall not acquire a red coloration on the addition of 1 drop of Phenolphthalein Solution. The *B.P.* includes no mention of the characters of the fatty acids obtained when an aqueous solution of the Soap is acidified with Diluted Sulphuric Acid, and the resulting fatty acids are filtered through a filter paper moistened with Water, washed till free from mineral acids and dried. They should possess the m.p. of about 45° C (113° F), an Iodine absorption of about 40 to 45 p.c., and a combining weight of about 278 to 280.

The more generally occurring impurities are the presence of an excess of alkali Hydroxide, an excessive amount of alkali carbonate, unsaponified oil or fat, Potassium Soap or excess of moisture. The behaviour of the filtered boiling Alcoholic Solution towards Phenolphthalein Solution and the titration of the hot aqueous solution of the residue left on the filter ensures the absence of free alkali Hydroxide, or an excessive amount of alkali Carbonate. Unsaponified oil or fat, if present, may be detected by a greasy stain being imparted by the Soap to white unglazed paper. The de quiescent nature of the ash remaining on ignition indicates the presence of Potassium Soap. It is officially required to lose when dried at a temperature of 110° C (230° F) about 30 p.c. of moisture. The *P.G.* requires that Hydrogen Sulphide Solution shall produce no change in a solution of 1 gramme of the Soap in 5 c.c. of Alcohol (90 p.c.)

SAPO DURUS.

HARD SOAP

FR, SAVON MEDICINAL, GER, MEDIZINISCHE SEIFE, ITAL, SAPONE
MEDICINALE, SPAN, JABON DE ACEITE DE OLIVAS

A solid, answering to the description given under 'Sapo Animalis,' but made by saponifying Olive Oil with Sodium Hydroxide. It is officially permitted to contain about 30 p.c. of Water.

Sapo Animalis is essentially Sodium Stearate, and Sapo Durus is essentially Sodium Oleate, but no confirmatory tests appear in the Pharmacopœia.

Solubility.—The greater part is soluble 1 in 20 of Water, entirely 1 in 1½ of boiling Water, 1 in 2 of boiling Alcohol (90 p.c.)

30 grains of White Castile Soap digested for 4 days in 1 oz. of cold Alcohol (90 p.c.), only 24 grains were dissolved, when heated it all dissolved.

Medicinal Properties.—Laxative and antacid. Combined with Rhubarb, it is administered as an antacid in dyspepsia attended with constipation. Large and frequent doses are most effective in removing gall-stones. Hard Soap, but more frequently Soft Soap, is made into a solution with warm Water, for use as an enema, ½ to 1 oz. to a pint.

The **Liniment**, which is made with soft soap, is used as a counter-

irritant, and is useful in sprains and rheumatic pains, and stiffness of joints

Dose —5 to 15 grains = 0.32 to 1 gramme

Prescribing Notes —*Best given in wafer paper or in cachets*

Official Preparations —Emplastrum Saponis, and Pilula Saponis Composita. Contained in Emplastrum Resine, Pilula Aloes Barbadosis, Pilula Aloes et Asafetide, Pilula Aloes Socotrine, Pilula Cambogae Composita, Pilula Rhei Composita, Pilula Scilla Composita. Used in the preparation of Hydragryll Oleas and Unguentum Zinci Oleatis. Soap Plaster is contained in Emplastrum Calcificans, and Emplastrum Cantharidis.

Not Official —Linimentum Saponis, Spiritus Saponatus, Eucaliol

Foreign Pharmacopœias —Official in Belg (Sapo Officinalis), Dan and Dutch (Sapo Medicatus), Hung (Sapo Venetus), Norw (Sapo Albus Oleaceus), Russ (Sapo Hispanicus Albus), Span (Jabon de Sosa), Swed (Sapo Medicatus), Swiss (Sapo Oleaceus), US (Sapo). With **Almond Oil** Fr (Savon Médicinal), Hung (Sapo Medicinalis), Ital (Sapone Medicinale), Mex (Jabon Medicinal), Port (Sabao Vegetal), Span (Jabon Amigdalino). With **Lard and Olive Oil**—Ger, Jap and Russ (Sapo Medicatus).

Tests —Hard Soap dissolves in Water, and readily in boiling Water and in boiling Alcohol. The *BP* requires that it shall not contain any free alkali Hydroxide as determined by digesting a weighed quantity of 5 grammes of the dried and powdered Soap in boiling Alcohol (90 p c), and whilst still hot filtering the solution through a filter and thoroughly washing it with boiling Alcohol (90 p c), the resultant filtrate should not produce a pink coloration with Phenolphthalein Solution. It also requires that it shall contain not more than 0.3 p c of alkali (Sodium) Carbonate as determined by titrating the residue from the above alcoholic solution with Tenth-normal Volumetric Sulphuric Acid Solution, not more than 3 c c of the solution should be required, 1 c c of Tenth-normal Volumetric Sulphuric Acid Solution is equivalent to 0.00526 gramme of Sodium Carbonate. The *USP* has a somewhat similar limit of alkalinity, but determines it by dissolving 5 grammes of the Soap in 50 c c of hot Water, and requires that when this solution is mixed with 3 c c of Tenth-normal Volumetric Oxalic Acid Solution the subsequent addition of a few drops of Phenolphthalein Solution should produce no pink or red tint. The alkali in combination of the fatty acids in the form of Soap may be determined as described under Sapo Animalis, by titrating the hot Alcoholic Solution used for the determination of the free alkali Hydroxide, with Tenth-normal Volumetric Sulphuric Acid Solution. The *USP* weighs the undissolved Sodium Carbonate from 20 grammes of Soap dissolved in Alcohol (94.9 p c), which should weigh not more than 0.8 gramme. Neither the *BP* nor the *USP* refers to the characters of the fatty acids obtained when an aqueous solution of the Soap is acidified with Diluted Sulphuric Acid. The liberated fatty acids filtered through a paper previously moistened with Water, washed till free from mineral acid and dried, should possess a m p of about 26° C (78.8° F), an Iodine absorption of about 80 p c and a combining weight of about 279.5.

The most common impurities are free alkali Hydroxide, excess of alkali, animal fats, and fatty acids from Oils other than Olive, Silica and other accidental impurities, unsaponified oil, Potassium soap and excess of moisture. Free alkali Hydroxide or excess of alkali Carbonate may be detected by the behaviour of the solution towards Phenolphthalein Solution and by the figure yielded on titrating the hot aqueous solution of the residue remaining after the filtration of the alcoholic solution as referred to at the commencement of the article. Animal fats may be determined by the gelation on cooling of the 25 p c solution of the Soap in Alcohol (94.9 p c), fatty acids from Oils other than Olive may be detected by determinations of the m p, Iodine absorption, and combining weight of the separated fatty acids. Metallic impurities may be detected by the Ammonium Sulphide and Hydrogen Sulphide test on the 1-20 Soap solution. Silica and other accidental impurities may be detected by a residue insoluble in Alcohol (94.9 p c) and in Water. Unsaponified oil leaves a greasy stain when the Soap is rubbed on white unglazed paper. Potassium Soap yields a deliquescent ash when the specimen is ignited with free access of air. The *BP* requires that the Soap shall lose, when dried at a temperature of 110°C (230°F), about 30 p c of moisture. In determining the amount of moisture the *USP* places 0.5 of a gramme of Soap with 10 c c of Alcohol in a tared beaker containing 1 gramme of clean dry sand, and evaporates to dryness drying the residue at 110°C (230°F) until of a constant weight. The *USP* requires that the loss should not exceed 36 p c.

Preparations

EMPLASTRUM SAPONIS. SOAP PLASTER

Hard Soap, 6, Lead Plaster, 36, Resin, 1. Melt each ingredient separately at a low temperature, mix, evaporate, with constant stirring, to a proper consistence (1 of Soap in $7\frac{1}{6}$).

New made with Hard Soap instead of Curd Soap

Emplastrum Saponis—*US*, 1 in 10, Emplastum Saponatum—*Aust*, about 1 in 14, *Dan*, 1 in 11, *Dutch* 1 in 10, *Ger* and *Jap*, about 1 in 17, *Hung*, about 1 in $15\frac{1}{2}$, *Norw*, about 1 in 17, *Russ*, 1 in $17\frac{1}{2}$, *Swiss*, 1 in 10, Emplastum Saponaceum—*Swed*, 1 in 9, Emplastro de Jabon—*Mex*, 1 in 18, Emplastro de Sabao—*Port*, 1 in $12\frac{1}{2}$, Emplastro de Jabon—*Span*, about 1 in 17, (Saponis Emplastum Camphoratum) *Belg*, Lead Plaster 75, Yellow Wax 10, Olei officinales 3, Camphor 2. *Aust* has also Emplastum Saponatum Salicylatum, Soap Plaster 85, Yellow Wax 5, Salicylic Acid 10.

LINIMENTUM SAPONIS. See *Sapo Mollis*

PILULA SAPONIS COMPOSITA. See *Opium*

Not Official

LINIMENTUM SAPONIS—Soap dried and granulated, 6, Camphor, in small pieces, 1, Oil of Rosemary, 1, Alcohol (95 p c), 72.5, Water, *q s* to make 100.

SPIRITUS SAPONATUS—Castile Soap, in shavings, 17.5; Alcohol (95 p c), 60, Water, *q s* to make 100. Dissolve and filter—*USNF*.

Spiritus Saponatus of the *P G* is made by saponifying Olive Oil 6 with solution of Potassium Hydroxide 7, and adding Alcohol 90, and Water 17, all by weight

Spiritus Saponatus of the *B P C* is made by dissolving 65 of Soft Soap in sufficient Alcohol (90 p c) to produce 100

EUNATROL (Sodium Oleate) Under this proprietary title has been introduced a substance containing pure Sodium Oleate. Stated to be useful as a cholagogue. Yellowish white fatty solid, possessing a faint odour of Oleic Acid. Best prescribed in pills or capsules. Dose, 10 to 15 grains = 0.65 to 1 gramme, twice daily. —*P J* '02, 16

SAPO MOLLIS

SOFT SOAP

A yellowish white, or yellowish green, unctuous semi solid

The *B P* Soft Soap is prepared with Potassium Hydroxide and Olive Oil. The *U S P* and the *P G* with Potassium Hydroxide, and Linseed Oil

Solubility — 1 in 4 of Water, 1 in 1 of boiling Water, almost entirely 1 in 1 of Alcohol (90 p c)

Official Preparation—**Limentum Saponis**. Contained in **Limentum Telesinthure**. **Soap Liment** is contained in **Limentum Opii**

Not Official—**Sapo Kalinus Venalis**, **Solutio Saponis Aetheracea**, **Spiritus Saponis Kalini Mollis**

Foreign Pharmacopœias Official in Austri, Belg, Dutch and Jap (**Sapo Kalinus**), Fr (**Savon Noir**) Ger, Hung, Russ, Swed and Swiss (**Sapo Kalinus** and **Sapo Kalinus Venalis**), Ital (**Sapone di Potassa**), U S (**Sapo Mollis**), Dutch has also **Sapo Superadipatus**, **Wool Fat 4**, **Soft Soap 20**, **Hard Soap 76**. Jap and Swiss have also **Sapo Venidis**

Tests—Soft Soap dissolves in cold Alcohol (90 p c), and readily in hot Alcohol (90 p c). It is officially required to yield no free alkali Hydroxide as determined by digesting 5 grammes in boiling Alcohol (90 p c), filtering and adding a few drops of Phenolphthalein Solution to the filtrate, which should not afford a red or a pink coloration. The *U S P* does not differentiate between the free alkali Hydroxide or the Carbonated alkali, but requires it to conform to the tests given below under the heading of Tenth normal Volumetric Oxalic Acid Solution. The *P G* requires that a solution of 10 grammes of Soap in 30 cc of Alcohol (90 p c) shall remain clear after the addition of 0.5 cc of Normal Volumetric Hydrochloric Acid Solution, and on the further addition of 1 drop of Phenolphthalein Solution shall not assume a red coloration. The *B P* fixes the limit of alkali Carbonate at 0.41 p c as determined by titrating the solution in hot Water of the residue insoluble in boiling Alcohol (90 p c). The use of Phenolphthalein Solution as an indicator of neutrality is recommended, but Methyl Orange Solution is more suited to the purpose, for the reason stated under **Sapo Durus**. Neither the *U S P* nor the *P G* includes a volumetric test for limit of alkali Carbonate. The *B P* fixes the limit of Potassium Carbonate, insoluble Soaps, etc., as determined by the weight of residue insoluble in hot Alcohol (90 p c), at 3 p c, the *U.S.P.* allows a similar limit for the amount of residue

insoluble in hot Alcohol (94.9 p.c.) No official mention is made as to the characters of the fatty acids, and consequently no confirmation is afforded that the particular Oil recommended in the official method of preparation has been employed in the manufacture of the Soap. To ascertain these characters the Soap is dissolved in Water, the aqueous solution acidified with Hydrochloric Acid, and the liberated fatty acids filtered through a filter paper previously moistened with Water, washed till free from mineral acid and carefully dried. They should possess a m.p. of about 26°C (78.8°F), an Iodine absorption of about 85 p.c., and a combining weight of 275 to 285. The *U.S.P.* does not include any methods for the examination of the fatty acids. The *P.G.* states that the fatty acid content of Sapo Kalinus Venalis amounts to at least 40 p.c. as determined by the process given in small type below. It has been recommended that a limit of Water should be added. The Soap may contain unsaponified oil or may be coloured with Copper salts, or be prepared by the saponification of Oils other than Olive. When rubbed on a piece of white glazed paper it should not impart an oily stain. When incinerated with free access of air it should yield an ash of a very deliquescent nature, which, when dissolved in diluted Hydrochloric Acid and tested with Hydrogen Sulphide, should not afford a brown coloration or precipitate, and which should impart a violet colour to a non-luminous flame. The presence of Oils other than Olive may be detected by the determination of the m.p., the Iodine absorption, and the combining weight of the fatty acids, obtained as above.

Alcohol—If 1 volume of a cooled solution of 5 grammes of Sapo Kalinus Venalis in 10 c.c. of hot Water be mixed with 1 volume of Alcohol, the mixture should remain clear, and even after the addition of 2 drops of Hydrochloric Acid a flocculent precipitate should not separate, *P.G.*

Tenth-normal Volumetric Oxalic Acid Solution—A solution of 5 grammes of Soft Soap in 50 c.c. of Water with 2 drops of Phenolphthalein T.S. added should require not less than 2.3 c.c. nor more than 4.5 c.c. of Tenth-normal Oxalic Acid Volumetric Solution to discharge the red tint, *U.S.P.*

Determination of the Fatty Acids—Dissolve 5 grammes of Soft Soap in 100 c.c. of hot Water. Mix the solution with 10 c.c. of diluted Sulphuric Acid in a test-glass and warm the mixture on a water-bath until the separated fatty acid forms a clear layer on the top of the aqueous fluid. Add 50 c.c. of Petroleum Benzine to the cooled liquid, stopper the test-glass and shake until solution of the fatty acid takes place, then allow 25 c.c. of this solution to evaporate at a gentle heat in a beaker and dry the residue to constant weight at a temperature not exceeding 75°C (167°F). The weight should be at least 1 gramme, *P.G.*

Preparation

LINIMENTUM SAPONIS. LINIMENT OF SOAP

Soft Soap, 2 oz., Camphor, 1 oz., Oil of Rosemary, 3 fl. oz., Alcohol (90 p.c.), 16 fl. oz., Distilled Water, 4 fl. oz. Dissolve the Soap in the Water, and mix it with the Camphor and Rosemary dissolved in the Alcohol, after a week, filter.

Tests—Soap Liniment has a sp. gr. of 0.895 to 0.900, it contains about 6 p.c. w/v of total solids and about 60 p.c. w/v of Absolute Alcohol.

This formula is practically the same as that which has been given in previous editions of the *Companion*, under the heading, 'Extractum Sarsæ Liquidum Compositum', the ingredients are similar to those of Decoctum Sarsæ Compositum, B P '85

Tests—Concentrated Compound Solution of Sarsaparilla has a sp gr of 1.020 to 1.040, it contains from 10 to 15 p c w/v of total solids and about 20 p c w/v of Absolute Alcohol

Not Official

DECOCTUM SARSÆ COMPOSITUM—Jamaica Sarsaparilla, cut transversely, 2½, Sassafras Root, in chips, ½, Guaiacum Wood turnings, ½, Dried Liquorice Root, bruised, ½, Mezerion Bark, ½, Boiling Distilled Water, 80. Digest the solid ingredients in the Water for an hour, boil for 10 minutes, cool, strain and make up to 20 fl oz.—B P 1885

This has been incorporated in the B P C

DECOCTUM ZITTMANNI FORTIUS Zittmann's Decoction (Strong)
—Sarsaparilla Root, 100, Water, 2600, Powdered Sugar, 6, Powdered Alum, 6, Mild Mercurous Chloride, 4, Red Mercuric Sulphide, 1, Anise Fruit, crushed, 4, Fennel Fruit, crushed, 4, Senna Leaves, cut, 24, Liquorice Root, cut, 12. The Sarsaparilla Root is digested for 24 hours with the Water, the powdered Sugar, powdered Alum, Mild Mercurous Chloride and Red Mercuric Sulphide are then added, the mixture heated in a covered vessel in a steam bath for 8 hours, stirring frequently, the Anise Fruit, Fennel Fruit, Senna Leaves and the Liquorice Root are added towards the end of the boiling, the liquid strained by expression and set aside for a short time. Decant 2500 parts of the clear liquid

DECOCTUM ZITTMANNI MITIUS Zittmann's Decoction (Weak) — The residue from the stronger decoction, and Sarsaparilla Root, cut, 50, Water, 2600, Lemon Peel, cut and bruised, 8, Cassia Bark, crushed, 8, small Cardamom Seeds, bruised, 8, Liquorice Root, cut and bruised, 8. The residue of the stronger decoction and the Sarsaparilla Root are extracted with Water by heating in a steam bath for 8 hours in a covered vessel, stirring frequently, the Lemon Peel, Cassia Bark, small Cardamoms and Liquorice Root are added towards the end of the operation, the liquid is strained by expression and set aside for a short time. Decant 2500 of the clear liquid

ROBERT'S DECOCTION—Sarsaparilla Root, in coarse powder, 1000, Water, q s. Place the Sarsaparilla Root in a closed vessel with 4000 of Distilled Water, and set aside for 8 hours, occasionally stirring, heat and keep boiling for 1 hour, then press out. Repeat this once. Evaporate the combined decoctions until there remains 1 litre (quart), mix well with an equal volume of Alcohol (90 p c), wash out the residue with boiling Alcohol (90 p c) ½ litre, strain through flannel and filter, evaporate to ½ litre or less, establish the quantity of Parillin and Sarsasaponin according to the method of V. Schulz (Christopherson), and adjust the strength of the finished product either by evaporating or by adding Distilled Water, so that it shall contain 2 p c of the above Glucosides

DECOCTUM SARSAPARILLÆ COMPOSITUM—Mix Sarsaparilla 20 with Water 520, and let the mixture stand for 24 hours at a temperature of 35° to 40° C, after the addition of Sugar 1, Potash Alum 1, heat in a covered vessel, stirring frequently, for 3 hours in a water bath. Add Anise 1, Fennel 1, Senna 5, Glycyrrhiza 2, leave in the water bath for a quarter of an hour and separate the liquid by pressing. After the pouring-off bring the weight of the decoction to 500 by the addition of Water.—Ger

TISANE DE SALSEPAREILLE—Macerate 50 grammes of Sarsaparilla (split and cut) in a little more than 1000 c c of Water for 2 hours, place it on the fire, and, as soon as it commences to boil, take it off again and let it digest for two hours, allow it to deposit and decant so as to obtain 1000 c c of Tisane.—Fr.

SYRUPUS SARSAPARILLÆ COMPOSITUS.—Fluid Extract of Sarsaparilla 2000, Fluid Extract of Glycyrrhiza 150, Fluid Extract of Senna 150,

Sugar 6500, Oil of Sassafras 2, Oil of Anise 2, Oil of Gaultheria 2, Water *q s* to make 10,000 — *U S*

Fluid Extract of Sassa-parilla 15, Fluid Extract of Glycyrrhiza 1, Fluid Extract of Senna 3, Spirit of Anise 1, — *Belg*

Percolate Sassa-parilla 100, Guaiac 100, and Anise Leaves 15, Sassafras 5, Anise 10, with a mixture of Alcohol (90 p c) and Water (equal weights) until 600 is obtained to 300 add 50 of Glycerin, filter and continue the evaporation to 100, to each 10 of this extract add 90 of Syrup — *Swiss*

SIROP DE SALSEPAREILLE COMPOSÉ — Pour on to 1000 of Sassa-parilla (split and cut in small pieces) to cover it, digest for 12 hours, separate the liquid from the residue in a similar manner and add the liquid to that of the first, also a third, which you put aside, heat this to ebullition and throw in 60 of dried Bojage flowers, 60 of dried petals of Rose, 60 of Senna leaves, and 60 of Anise fruits, allow to infuse for 6 hours and press, evaporate the first liquors, and the residue is reduced to 500 g, mix with the third liquid and continue the evaporation until the product weighs 1000 g, then clarify with white of egg and strain this on a cloth, add to the liquid thus obtained 1000 of Sugar and 1000 of White Honey and make a syrup by boiling and clarification until it has a sp gr of 1.29 — *Fr.*

SASSAFRAS RADIX.

SASSAFRAS ROOT.

The dried Root of *Sassafras officinale*, T. Nees and Eberm.

It contains a yellowish, or reddish yellow, volatile Oil (Oil of Sassafras), which is largely distilled in America and is official in U S, the yield is about 2 p c. The bulk of this Oil consists of *Safrol*, $C_{10}H_{16}O_2$, a compound also extracted from Oil of Camphor. It is much used for scenting soap.

Medicinal Properties — Aromatic and carminative. Used as an adjuvant to other medicines.

The oil strongly recommended for pediculi; the brush is dipped in a saucer full of the oil, the whole head well brushed with it, and a close-fitting linen cap put on for 24 hours — *B M J '07*, 1164.

Official Preparation — Contained in Liquor Sassa Compositus Concentratus.

Foreign Pharmacopœias — Official in Austri, Mex and Port, the Root, Ger, Jap, Span, Swiss and U S, the Root-bark.

Descriptive Notes — The root met with in commerce is usually offered by the wholesale houses in the form of chips, apparently of large roots, since very little bark is present. The bark is rough, brown, with a whitish external layer, but smooth, with a satiny lustre on the inner surface. It is slightly astringent and has a *Safrol* flavour. The chips of wood are greyish, with a yellowish or often with a reddish tinge. In the U S P the much more aromatic bark, deprived of the periderm, is official, as well as the pith of the stem (*Sassafras Medulla*) see below.

The bark is characterised by large oil cells, pitted parenchymatous cells, thick-walled bast fibres, and starch grains singly or in groups of 2 to 3 each, with a well-marked hilum. The starch also occurs in the wood and in the medullary rays. The bark contains a dark brown colouring matter soluble in Liquor Potassæ.

Tests.—Sassafras Root yields about 2 p c of ash

SASSAFRAS MEDULLA Sassafras Pith (*U S*)—It abounds in a gummy matter, which forms a mucilage with Water. 60 grains of Pith to 20 fl oz is used as a soothing application to the eyes, and as a drink in diarrhoea

OLIVERI CORTEX *Syn* BLACK SASSAFRAS—The dried Bark of *Ornamomum Oliveri* is official in the *Ind* and *Col Add* for the Australian Colonies. Also *Tinctura Oliveri Corticis*, 1 in 10 (Alcohol 60 p c), dose, 30 to 60 minims = 1 3 to 3 6 c c

SCAMMONIÆ RADIX.

SCAMMONY ROOT

The dried brownish, or yellowish-grey, perennial tapering Root of *Convolvulus Scammonia*, L

From Syria and Asia Minor

Official Preparation.—Used in the preparation of Scammonia Resina

Official in Spain and Swiss

Descriptive Notes—The root occurs in commerce in stout hard, cylindrical pieces, often spirally twisted, and having a rough, furrowed, greyish-brown bark, and it is often 2 to 3 in (50 to 75 mm) in diameter. Internally it is greyish-yellow and fibrous, and in transverse section exhibits irregularly arranged circles of woody bundles. In the cortical region and around these circles dark resin cells are frequent, and the softer tissues are full of a mullet-shaped starch, which is characteristic. The root has been scarce during late years, its exportation having been prohibited, and a root known in commerce as 'Mexican Scammony' Root, derived from *Ipomoea Orizabensis*, Leden, has taken its place as a cheap substitute for the manufacture of Scammony Resin, with which its resin appears to agree in chemical characters. It occurs in irregularly oblong segments, evidently derived from a large root, and like Scammony Root has projecting fibres, but differs in its radiate structure. It contains about 15½ to 18½ p c of resin against 5½ to 8½ p c in true Scammony Root. See *P J* (4) xviii pp 326, 327

Tests—Scammony Root yields about 10 p c of ash. It is officially stated to yield a resin possessing the properties of Scammony Resin when treated with Alcohol (90 p c), but no indication as to the amount expected to be yielded is given. It yields about 9, if it is of Levantine origin, the average yield being about 8 p c. As the root is used only for preparing the Resin it is considered that probably no standard need be indicated, consideration should, however, be given to the different varieties of Scammony Resin now being obtained from roots of the so-called Mexican Scammony

SCAMMONIÆ RESINA.**SCAMMONY RESIN**

Greenish-grey, or brownish-green, translucent, brittle lumps, with more or less sharp edges, and breaking with a shining fracture. It has a peculiar, characteristic odour.

Scammony Resin is official in the *BP* and the *USP*, but not in the *PG*. It is identical with the Ether-soluble Resin of Jalap.

It is prepared by extracting Scammony Root with Alcohol (90 p c) recovering the greater part of the Alcohol and pouring the concentrated liquid into Distilled Water, which precipitates the Resin.

Solubility.—It is soluble in almost all proportions of Alcohol (90 p c) or Ether, also soluble in Solution of Potassium Hydroxide.

The purified Resin is known in this country as **Scammonin**, see p. 705.

Medicinal Properties.—An energetic, hydragogue cathartic. May be used when brisk action is needed, as in cerebral congestion and severe dropsy, but on account of its griping properties it is rarely used alone. In combination it promotes the action of other medicines, whilst its own harshness is mitigated. It acts also as an anthelmintic, to round-worms and tapeworms.

Dose.—3 to 8 grains = 0.2 to 0.52 gramme.

Official Preparations.—*Pilula Scammonii Composita* and *Pulvis Scammonii Compositus*. Contained in *Extractum Colocynthis Compositum*, *Pilula Colocynthis Composita*, and *Pilula Colocynthis et Hyoseyami*.

Not Official.—*Confectio Scammonii*, *Mistura Scammonii*, *Pulvis Scammonii cum Hydrargyro*.

Foreign Pharmacopœias.—Official in Belg., Fr., Ital., Mex., Norw. and U.S.

Tests.—Over and above the official description of the Resin the *BP* does not give any chemical tests or constants by which Scammony Resin may be distinguished. The Acid Value and Saponification values afford a means of distinguishing the Resin. The Acid value should be, according to Kremel, 14.6, the Ester value 171.0 and the Saponification value 185.6. It may be distinguished from Guaiacum Resin by the non-production of a blue coloration when Ferric Chloride T.S. is added to its solution in Absolute Alcohol, and by the non-formation of a blue colour on the addition of Hydrogen Peroxide Solution to its solution in Absolute Alcohol. It may be distinguished from Jalap Resin by the fact that it dissolves almost entirely in Ether. It should yield when incinerated with free access of air not more than 1 p c of ash, which is also the limit allowed by the *USP*. The *BP* gives no figure for the ash limit. Resin, if present, may be detected by the increase in the Acid value and the decrease in the Ester value.

Preparations

PILULA SCAMMONII COMPOSITA. COMPOUND SCAMMONY PILL

Scammony Resin, 1, Jalap Resin, 1; Curd Soap, in powder, 1, Tincture of Ginger, 3, dissolve, and evaporate to pill consistence.

Dose—4 to 8 grains = 0.26 to 0.52 gramme.

PULVIS SCAMMONII COMPOSITUS COMPOUND POWDER OF SCAMMONY

Scammony Resin, 4, Jalap, 3, Ginger, 1 (1 in 2)

Dose—10 to 20 grains = 0.65 to 1.3 grammes

Not Official

CONFECTIO SCAMMONII—Resin of Scammony, in powder, 6, Ginger, 3, Oil of Caraway, $\frac{1}{2}$, Oil of Cloves, $\frac{1}{2}$, Syrup, 6, Clarified Honey, 8 Rub the powder with the Syrup and Honey, then add the Oils, and mix. **Dose**—10 to 30 grains—*B P* 1885

This has been incorporated in the *B P C*

MISTURA SCAMMONII—Scammony, in powder, 6 grains, Milk, 2 fl oz—*B P* 1885

This has been incorporated in the *B P C*

PULVIS SCAMMONII CUM HYDRARGYRO—Mercurous Chloride, 1, Scammony Resin, in powder, 4—*St Thomas's*

This has been incorporated in the *B P C*

Tabellæ Scammoniae cum Chocolata official in Belg., about 3 grains in each

SCAMMONIUM.

SCAMMONY

Brown, dark grey, or brownish-black, irregular masses, or circular cakes, breaking with a glossy, resinous fracture. It possesses a peculiar, cheese-like odour.

It is officially described as a Gum-resin, obtained by incision from the living Root of *Convolvulus Scammonia*, L., known in commerce as Virgin Scammony.

Chiefly from Smyrna, in Asia Minor.

Solubility—Almost entirely dissolved in boiling diluted Alcohol.

Medicinal Properties.—Similar to those of Scammony Resin, but Scammony emulsifies with Water, the Resin does not.

Dose—5 to 10 grains = 0.32 to 0.65 gramme.

Foreign Pharmacopœias—Official in Fr., Mex., Port., Span. and U.S.

Descriptive Notes—Scammony Resin is usually imported in boxes containing pieces varying in purity, which are sorted in this country, the purest pieces being sold as Virgin Scammony. The finest pieces are brittle, translucent, have a brownish tint, a resinous fracture, and are more or less covered with a greyish powder; other pieces have a blackish resinous fracture, are less brittle, and sometimes more or less porous, due to fermentation after collection. An inferior kind, adulterated with Flour and Chalk, known as Skilip Scammony, is hard, tough, not easily fractured, and has an opaque, greyish fracture; it contains only about 40 p.c. of Resin. When rubbed with a wetted finger Scammony gives a milky emulsion, which Resin of Scammony does not. Formerly, Aleppo Scammony was purer than that of Smyrna, now the reverse is the case.

Tests.—Scammony is readily friable and forms with Water a greenish emulsion. When treated with Ether at least 70 p.c. of Resin is officially required to be dissolved. In estimating the Resin soluble in Ether it is recommended to use a light Ether, sp. gr. 0.717, and to break up the residue after evaporating the Ether and again heat, in order to avoid error due to the Resin holding down the Ether. It would probably be better to dry the Scammony, extract with Ether and weigh the insoluble residue. The *U.S.P.* requires that not less than 75 p.c. should be soluble in Ether, and this was the standard previously adopted in the *B.P.* 1885. It may be noted that the majority of good commercial samples of the Gum Resin yield a larger percentage of Ether-soluble Resin than the present official limit. The *U.S.P.* adds that the residue remaining after the evaporation of the ethereal solution when dissolved in hot Water, Sodium Hydroxide Solution is not reprecipitated on acidification with Diluted Sulphuric Acid. Aleppo Scammonium has an Acid value of 8.2, an Ester value of 172.0 and a Saponification value of 180.2 (Dieterich, *Analysis of Resins, Balsams and Gum Resins*). It is generally heavily adulterated, Resin, Starch, Chalk and other mineral substances being added. Resin if present, may be detected by the increase in the Acid value and the decrease in the Ester value. A portion of the powdered Gum Resin when boiled with Water and cooled should give no decided blue coloration on the addition of Ferric Chloride Solution. The greenish emulsion formed on triturating the powdered Gum Resin with Water should not effervesce on the addition of Diluted Hydrochloric Acid. It should leave when ignited with free access of air not more than 3 p.c. of ash which is the official limit; the *U.S.P.* also places the ash limit at this figure. It may be distinguished from Guaiacum Resin by not affording a blue coloration on the addition of Ferric Chloride T.S. to its alcoholic solution.

SCILLA.

SQUILL

FR., SCILLE, GLR., MITTWEIHERL., ITAL., SCILLA; SPAN., ESCILA

The Bulb of *Urginea Scilla*, Steud., divested of its dry membranous outer scale, cut into slices and dried.

From the Mediterranean coast.

Two active principles have been extracted from Squill, Scillitoxin (Scillanin) and Scillipicrin, the former strongly affects the heart, but their actions are antagonistic.

Under the title of *Urginea*, *syn.* Indian Squill, the younger bulbs of *Urginea Indica*, Kunth, also the younger bulb of *Scilla Indica*, J. J. Baker, are official in the *Ind.* and *Col.* Add for India and the Eastern Colonies.

Medicinal Properties.—A stimulant expectorant, diuretic and cardiac tonic, acting similarly to Digitalis, but is more irritating to the gastro-intestinal tract than Digitalis. It increases the secretion of the bronchial mucous membrane, and the expectoration of mucus. In chronic bronchitis it is used in combination with other

expectorants, such as Ipecacuanha and Ammonia. In acute bronchitis it is too irritating to the bronchial mucous membrane, while in phthisis it may produce dyspepsia. In dropsy, especially if cardiac in origin, it is combined with Blue Pill and Digitalis.

Dose—1 to 3 grains = 0.065 to 0.2 gramme

Official Preparations—Acetum Scillæ, Oximel Scillæ, Pilula Scillæ Composita, and Tinctura Scillæ. Contained in Pilula Ipecacuanhæ cum Scilla. The Vinegar is used in the preparation of Sympus Scillæ.

Not Official—Sympus Scillæ Compositus, Fluidextractum Scillæ, Mistura Scillæ Composita, Mistura Scillæ et Ipecacuanhæ, Linctus Scillæ, Linctus Scillæ Opiatus, Dr. Abercrombie's Cough Mixture, Dr. Milne's Fotheigill's Mixture.

Foreign Pharmacopœias—Official in all the Foreign Pharmacopœias, Fr. (Scille), Mex. and Span. (Escila) Belg. and Fr. have Extractum Scillæ, and U.S. has Fluidextractum Scillæ.

Descriptive Notes—The Squill bulbs, which are often 6 in. (15 cm.) in diameter, are offered in commerce in the form of small curved dried strips about 1 to 5 cm. long, and $\frac{3}{8}$ to $\frac{1}{2}$ in. (9 to 15 mm.) broad in the middle, tapering to either end, usually of a yellowish-white colour, tough and slightly flexible, but brittle when quite recently dried. It has no odour, but a disagreeable bitter taste. There are two varieties of the bulb, known respectively as the Red and White Squill. When derived from the Red Squill the strips have a pinkish colour. Occasionally an unusually bitter sample is met with, but the cause of this has not been ascertained. Squills are very hygroscopic, and to keep their medicinal activity unimpaired should be thoroughly dried on arrival, and kept in an air-tight vessel. The powder is best kept in a bottle with a hollow stopper containing quicklime, or it readily cakes into a hard mass. The bulbs are collected in August, and when fresh their handling causes considerable irritation to the skin. Squill is characterised by the presence of long prismatic crystals of Calcium Oxalate, often 1 mm. long, immersed in a mucilage which contracts into a jelly on the addition of Alcohol. Acicular raphides are also present. Large stomata also occur, and small bundles of laticiferous vessels. Starch granules in small quantity are present in elongated cells near the vascular bundles.

Tests—Squill yields from 2 to 3 p.c. of ash, and $\frac{1}{2}$ p.c. should not be exceeded. Determinations of the ash made in the author's laboratory showed an average of 2.4 p.c. A standard of 20 p.c. has been suggested for the amount of moisture.

Preparations

ACETUM SCILLÆ VINEGAR OF SQUILL

$2\frac{1}{2}$ of Squill, bruised, macerated with Diluted Acetic Acid, $7\frac{1}{2}$ to yield 20 (1 in 8)

It is conveniently filtered through Talc.

Dose—10 to 30 minims = 0.6 to 1.8 c.c.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Mex., Norw., Port., Swiss and U.S., 1 in 10. All by weight except U.S.

Tests.—Vinegar of Squill has a sp gr of 1.035 to 1.040, it contains from 5 to 9 p c w/v of total solids and about 4.0 p c w/v of absolute Acetic Acid, as determined by titrating a measured quantity (10 c c.) with Normal Volumetric Sodium Hydroxide Solution, 10 c c requiring from 6 to 7 c c.

Although a standard of 9 p c w/v of total solids is sometimes reached, it appears (*C D* '02, 1733, 808) that a fair average is 8 p c, with 7 to 9 p c as the limit, the figures outside by a fraction. When prepared strictly in accordance with the *B.P.* directions it contains immediately when made 3.6 to 4.0 p c of absolute Acetic Acid. The *German Pharmacopœia* allows for a loss of Acetic Acid in the process, as the menstruum contains only 5.4 p c w/w of Acetic Acid, but the titration test of that *Pharmacopœia* requires 4.8 to 5.1 p c w/v of the acid in the finished Vinegar.

A corresponding preparation, *Acetum Urgineæ*, is official in the *Ind* and *Col. Add* for India and the Eastern Colonies.

OXYMEL SCILLÆ. OXYMEL OF SQUILL

Squill, bruised, 2½, Acetic Acid, 2½, Distilled Water, 8, Clarified Honey, liquefied, q s to bring the fluid to sp gr 1.320
(1000 + 1 in 15)

Dose—½ to 1 fl drm = 1.8 to 3.6 c c

A corresponding preparation, *Oxymel Urgineæ*, is official in the *Ind* and *Col. Add* for India and the Eastern Colonies.

Foreign Pharmacopœias—Official in Austr, Extract of Squill 1, Acetic Acid (96 p c) 1, Refined Honey 98, Dan and Norw, Vinegar of Squill 35, Honey to make 100, Dutch, Vinegar of Squill 2, Refined Honey 1, Sugar 1, Ger Ital, Jap and Span, Vinegar of Squill, 1 Refined Honey 2, Hung, Refined Honey 320, Extract of Squill 2, Acetic Acid (96 p c) 3, Diluted Acetic Acid 4, Port, Mex and Gr, Vinegar of Squill 1, Honey 4, Swed, Squill 1, Acetic Acid 2, Distilled Water 8, Refined Honey 30, Swiss, Vinegar of Squill 3, Sugar 3, Refined Honey 4. Not in Belg, Russ or U S.

Tests—Oxymel of Squill should have a sp gr of about 1.320

PILULA SCILLÆ COMPOSITA. COMPOUND SQUILL PILL

Squill, 1½, Ginger, 1, Ammoniacum, 1 Hard Soap, 1, Syrup of Glucose (by weight), about 1
(about 1 in 4)

Dose—4 to 8 grains = 0.26 to 0.52 gramme

A corresponding preparation, *Pilula Urgineæ Composita*, is official in the *Ind* and *Col. Add* for India and the Eastern Colonies.

SYRUPUS SCILLÆ. SYRUP OF SQUILL

Vinegar of Squill, 20, Refined Sugar, 38, it should yield 58 by weight
(about 1 of Squill in 18)

Quantity of Sugar reduced from 40 to 38

Dose.—½ to 1 fl drm = 1.8 to 3.6 c c

Official in U S, Vinegar of Squill 45, Sugar 80, Water to measure 100

Syrupus Scillæ Compositus—Fluid Extract of Squill, 8, Fluid Extract of Senega, 8, Antimony and Potassium Tartrate, 0.2, Purified Talc, 2, Sugar 75, Water, q s to make 100. Average Dose—2 p c (30 minims).

A corresponding preparation, *Syrupus Urgineæ*, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Tests—Syrup of Squill has a sp gr of about 1.345

TINCTURA SCILLÆ TINCTURE OF SQUILL

1 of Squill, bruised, macerated with 5 of Alcohol (60 p c) (1 in 5)

Dose—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Belg., Fr., Ger., Ital., Jap., Mex., Port., Span. and Swiss, 1 in 5, all by weight, U.S., 1 in 10. Fr. has also *Vin de Scille Composé*.

A corresponding preparation, *Tinctura Urgineæ*, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Tests—Tincture of Squill has a sp gr of 0.960 to 0.975, it contains about 12 p c w/v of total solids and about 54 p c w/v of Absolute Alcohol. A standard of 10 p c w/v or more has been suggested for the total solids

Not Official

FLUIDEXTRACTUM SCILLÆ—Squill, in No. 20 powder, 100, Acetic Acid and Water q s. Mix 27.5 of Acetic Acid with 72.5 of Water and macerate the powder in 80 of the mixture for 48 hours, transfer to a percolator and by slow percolation with the same menstruum continue until the product measures 100—*U.S.P.* Average Dose—1½ minims (0.1 c c)

This is an Acetic Acid extract

The previous edition *U.S.P.* 1890 was prepared by exhausting 100 of Squill with Alcohol (70 p c), reserving the first 75 and evaporating the weaker percolates to an extract, which was dissolved in the reserved portion, and the product made up to 100. The *B.P.C.* have included an *Extractum Scillæ Liquidum* made in a similar manner to this (*U.S.P.* 1890) with Alcohol (70 p c). It has been proposed by Greenish (*P.J.* '07, ii 99) to make *Acetum Scillæ*, *Oxymel Scillæ*, *Syrupus Scillæ*, with this fluid extract, but in that case the solvent action of the Acetic Acid would be lost. The *U.S.P.* have now discarded this fluid extract for one made with Acetic Acid as given above.

MISTURA SCILLÆ COMPOSITA—Oxymel of Squill, 40 minims, Compound Tincture of Camphor, 20 minims, Spirit of Nitrous Ether, 20 minims, Water, to 1 fl oz.—*East London*

This has been incorporated in the *B.P.C.*

MISTURA SCILLÆ ET IPECACUANHÆ—Vinegar of Squill, 10 minims, Vinegar of Ipecacuanha, 10 minims, Potassium Citrate, 15 grams, Solution of Ammonium Acetate, 2 fl dm, Anise Water, to 1 fl oz.—*St. Thomas's*

This has been incorporated in the *B.P.C.*

LINCTUS—Oxymel of Squill, 15 minims, Mucilage of Tragacanth, 15 minims, Glycerin, 15 minims, Emulsion of Chloroform, 3 minims, Syrup, to 1 fl dm.—*St. Thomas's*

This has been incorporated in the *B.P.C.* under the title *Linctus Scillæ*, *syn* Linctus, Simple Linctus

LINCTUS SCILLÆ OPIATUS—Compound Tincture of Camphor, Oxymel of Squill, Syrup of Tolu, of each equal parts. Dose—One teaspoonful—*St. Bartholomew's*

This has been incorporated in the *B.P.C.* under the title *Linctus Scillæ Compositus* (or *Opiatus*)

DR. ABERCOMBIE'S COUGH MIXTURE—Tincture of Opium, 160 minims, Syrup of Squill, 2 fl oz, Cinnamon Water, 4 fl oz, Water, 4 fl oz. One tablespoonful for a dose.—*Pharm. Form.*

This has been incorporated in the *B.P.C.* under the title *Mistura Scillæ et Opii*, *syn* Abercombe's Cough Mixture

DR MILNER FOTHERGILL'S MIXTURE—Syrup of Squil, 1, Dilute Hydrobromic Acid, $\frac{1}{2}$, Spirit of Chloroform, $\frac{1}{2}$, Water to 8 Dose $\frac{1}{2}$ or 3 times a day, to be sipped slowly — *Pharm Form*

This has been incorporated in the *B P C* under the title *Mistura Scillæ* with the synonym, Fothergill's Cough Mixture, but with Water to 10 instead of to 8

SCOPARII CACUMINA.

BROOM TOPS

The fresh and the dried Tops of *Cytisus scoparius*, Link, a woody shrub indigenous to England, and also found throughout the greater part of Europe

Medicinal Properties—Diuretic and in large doses cathartic. Employed in dropsical complaints, especially if cardiac, and often prescribed along with Potassium salts and Digitalis, in renal dropsy it is contra-indicated if there be acute nephritis

Official Preparations—Infusum Scoparii and Succus Scoparii

Not Official—Decoctum Scoparii, Infusum Scoparii Concentratum, Sparteina, Sparteinae Pericardium, Sparteinae Sulphas

Foreign Pharmacopœias—Official in U S (dried tops)

Descriptive Notes—Both the fresh and dried tops are official, the former for the succus and the latter for the infusion. The wiry dark green stem and branches are 5-angled, hairy on the young twigs (glabrous, *B P*), flexible, and $\frac{1}{2}$ to $\frac{1}{3}$ in (2 to 3 mm) thick. The branches are set at an acute angle to the stem, and in the upper part have simple and nearly sessile leaves, but in the lower part trifoliate leaves, which are obovate and are furnished with a petiole about their own length. The fresh plant, when bruised, has a characteristic odour, which is lost in drying. It has a faintly bitter taste. The younger parts of the plant are hairy as well as the young leaves, and the young branches are pubescent before the plant flowers, but in the dried drug (probably collected after the fruit is formed) the pubescence of the stem and the hairs of the leaves are not usually visible, and the *B P* description evidently applies to the plant collected in summer or early autumn.

Tests Broom Tops yield about 3 p.c. of ash

Preparations

INFUSUM SCOPARII INFUSION OF BROOM

Broom Tops, dried and bruised, 2, boiling Distilled Water, 20; infuse 15 minutes, and then strain (1 in 10)

Dose.—1 to 2 fl oz = 28.4 to 56.8 c.c.

It takes the place of Decoction of Broom, *B P* '85

SUCCUS SCOPARII, JUICE OF BROOM

To 3 of Juice from bruised Fresh Broom Tops, add 1 of Alcohol (90 p.c.), after 7 days filter.

Dose.—1 to 2 fl drms = 3.6 to 7.1 c.c.

Not Official

DECOCTUM SCOPARII—Broom Tops, dried, 1, Distilled Water, *qs* to make the final product after boiling for 10 minutes measure 20 fl oz.—*B P* 1895
This has been incorporated in the *D P C*

INFUSUM SCOPARII CONCENTRATUM Broom Tops, in No 20 powder 80, Alcohol (90 p c), 25, Dilute Chloroform Water (1 in 1000), *qs* to make 100 Prepare by reprecipitation Before the addition of the Alcohol to the reserved portion this should be heated to a temperature of not less than 85° C and maintained therefor 5 minutes Dose—1 to 2 fl drim = 5 to 7 1 —*Flan*, and *Wright, P J* '06, 1 165 and '07, 1 621, *C D* '06, 1 252, and *Y B P* 1907, 348

This appears in the *D P C*

SPARTEINA ($C_{11}H_{16}N$, eq 232.53)—A clear, colourless, oily liquid, heavier than Water, having an odour somewhat resembling Aniline, and an intensely bitter taste It is a liquid alkaloid, obtained from Broom

It should be kept in well stoppered glass bottles of a dark amber tint, and protected as far as possible from exposure to light and air, as it tends to darken in colour and to become thick.

Practically insoluble in Water, soluble in Alcohol, in Ether, and in Chloroform

Foreign Pharmacopœias—Official in Mex and Spun

Tests.—Sparteine boils at about 287° C (548.6° F) It dissolves in Alcohol (90 p c), the solution being levogyrate It possesses a strongly alkaline reaction towards the usual indicators of neutrality A glass rod moistened with Hydrochloric Acid held over a watch glass containing a drop of Sparteine evolves white fumes It unites with acids to form crystallizable salts On gradually adding a solution containing 3 parts of Iodine dissolved in Ether to an ethereal solution of 1 part of Sparteine, a black precipitate is formed, which when separated, washed with Ether and dissolved in boiling Alcohol crystallises on cooling in beautiful green needles A solution of Sparteine gives with Cadmium Iodide Solution a white curdy precipitate, with Sodium Phosphomolybdate Solution a white precipitate, redissolving on heating the liquid Platinum Chloride Solution yields a yellow crystalline precipitate very insoluble in cold Water and Alcohol, but crystallising from Hydrochloric Acid in rhombic prisms It yields no coloration with Sulphuric or Nitric Acid It may be quantitatively determined by titration with Normal Volumetric Sulphuric Acid Solution, using Iodocarm Solution as an indicator of neutrality In a purely aqueous titration 1 c.c. of Normal Volumetric Sulphuric Acid Solution represents 0.11626 gramme of pure Sparteine In strong alcoholic solution (provided the Water introduced is not sufficient to unduly reduce the alcoholic strength), 1 c.c. of the Normal Acid Solution represents 0.23253 gramme of pure Sparteine It should leave no weighable residue when heated with free access of air.

SPARTEINÆ PERIODIDUM ($C_{11}H_{16}N$ 2HI eq 864.04)—Small, bronze green crystals, or bronze green amorphous powder Insoluble in Water, soluble in Alcohol (90 p c) Diuretic Prepared by the author for the late Dr Mortimer Granville, and forming one of the series of alkaloid periodides which evolve Iodine slowly, and which were used by him in gout

Dose— $\frac{1}{2}$ to 1 grains = 0.032 to 0.26 gramme

Tests.—Sparteine Periodide dissolves in Alcohol (90 p c) 1 gramme when dissolved in 10 c.c. of Alcohol (90 p c) and titrated with Tenth normal Volumetric Sodium Thiosulphate Solution requires about 33.0 c.c. to discharge the colour produced on the addition of Starch Mucilage This indicates about 11.5 p.c. of Iodine When treated with Sodium Thiosulphate Solution and made alkaline with Ammonia, shaken with Ether and the ethereal solution spontaneously evaporated, the separated alkaloid should answer the tests given under Sparteine When ignited with free access of air it should leave no weighable residue

SPARTEINÆ SULPHAS ($C_{11}H_{16}N_2 \cdot H_2SO_4 \cdot 5H_2O$, eq 419.27)—Colourless, prismatic crystals, or a white crystalline powder

The number of molecules of Water of crystallisation varies with the solution from which it is crystallised The *U S P* states that when recrystallised from a

solution in Alcohol (48 p c) to contain 5 molecules of Water of crystallisation. The salt official in the *F. C.* (1905), and the salt official in the present *U S P*, contains 5 molecules of Water of crystallisation, that official in the 1890 edition of the *U S P* contained 4 molecules of Water of crystallisation.

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from the light.

Soluble 2 in 1 of Water, 1 in 5 of Alcohol (90 p c).

Medicinal Properties—Cardiac tonic and diuretic. Useful in mitral disease. It slows and strengthens the pulse. Its action is persistent than that of Digitalis.—*B M J* '86, 1 1246, '88, 1 *P J* (3) 211 543, *P* 1 213, as a preliminary to chloroform.—*B M J E* '94, 11 48, *T G* '95, 40.

Dose— $\frac{1}{2}$ to 2 grains = 0.016 to 0.13 gramme.

Foreign Pharmacopœias—Official in *F.*, *Mex.*, *Span.*, *Swiss* and *U.S.*

Tests—Sparteine Sulphate when heated to a temperature of 110° C (230° F) loses its Water of crystallisation, equivalent to 21.3 p c. The anhydrous salt melts at 136° to 138° C (276.8° to 280.4° F), the *F.* *Codex* gives 145° C (293° F). It dissolves readily in Water, forming a clear solution possessing in acid reaction towards blue Litmus paper. When rendered alkaline with Ammonia, shaken with Ether and the ethereal solution evaporated spontaneously, a solid answers the tests given under Sparteine. A 5 p c solution of the salt in Water gives a yellow precipitate with Potassium Ferrocyanide Solution. A small quantity of the salt mixed in a porcelain capsule with one-third of its weight of Chromic Acid, and gently warmed gives a green coloration and simultaneously emits a distinct odour of Camphor. An aqueous solution of the salt mixed with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. The percentage of Sparteine Sulphate present may be determined by direct titration with Tenth-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator. The choice of a solvent in performing the process of titration is important, as although Sparteine is di-basic only half the acid is indicated by titration under the above conditions in aqueous solutions, but when Sparteine Sulphate is dissolved in Absolute Alcohol (90 p c) the full quantity of acid is indicated. When dissolved in Water 1 c c of Tenth-normal Volumetric Sodium Hydroxide Solution is equivalent to 0.041927 gramme of crystallised Sparteine Sulphate. When dissolved in Absolute Alcohol (90 p c) as a solvent, each c c of the solution corresponds to 0.02096 gramme of the pure crystallised salt free from readily charred organic impurities, and should not contain Ammonium salts, or mineral matter. The salt itself and its solution in concentrated Sulphuric Acid should be colourless. It should yield no odour of Isophenylcyanide when 1 dgm is heated with 5 drops of Chloroform and 1 c c of Alcoholic Potassium Hydroxide Solution, indicating the absence of Aniline Sulphate. It should yield no ammonia-like odour when warmed with Potassium Hydroxide Solution, indicating the absence of Ammonium salts, although a piece of red Litmus paper suspended in the mouth of the test-tube will gradually acquire a blue colour. When mixed with free access of air the salt should leave no weighable residue, indicating the absence of mineral impurities.

Hypodermic Lamels, $\frac{1}{2}$ grain of Sparteine Sulphate in each.

Oxysparteinae and **Oxysparteinae Hydrochloridum** and **Sulphas** have been used in medicine, the dose being about the same as that of Sparteina.

Not Official **SCOPOLA.**

The dried Rhizome of *Scopola Carmichaelia*, Jacq., known also on the Continent as *Scopolia atropoides*, Link.

The dried Rhizome is official in the *U S P*, and is required to yield not less than 0.5 p c of mydriatic alkaloids,

It contains Hyoscyamine, Scopolamine (amorphous Hyoscine) and Atroscine (crystalline Hyoscine)

Medicinal Properties—It has the same properties as Belladonna and Hyoscyamus

This drug has not 'taken' in British practice, but it is used on an immense scale in America for the preparation of what is termed 'Belladonna' plaster

Action of Scopolamine Hydrochloride on the eye—*P*, liv 469, *TG* '93, 338, 781, '94, 423, 480, 625, 680, *BMJ* '94, ii 497

Foreign Pharmacopœias—Official in Jap and U S Jap has an **extract** prepared with weak Alcohol, a **plaster**, Extract 1, Resin Plaster 9, a **tincture**, Root 1, Dilute Alcohol 5, and an **ointment**, Extract 1, Laid 9 U S has extract and fluid extract (*see below*)

Tests—10 grammes of the dried Rhizome in No 60 powder when examined by the *USP* process for assay of Belladonna Leaves, given under Belladonna Folia, should yield a quantity of mydriatic alkaloids corresponding to not less than 0.5 p c

EXTRACT SCOPOLA—The Extract of Scopola (*USP*) is prepared by evaporating the Fluid Extract to a pulular consistence in a porcelain dish at a temperature not exceeding 50° C (122° F), constantly stirring during the evaporation. It is required to contain 2 p c of mydriatic alkaloids, and the *USP* directs that should the Extract be found to contain more than this percentage sufficient powdered Milk Sugar should be added to reduce it to this standard

Tests—The *USP* method of assay is identical with that described under the *USP* process for the assay of Extract of Belladonna Leaves described under Belladonna Folia. Inasmuch, however, as 2 grammes of the Scopola Extract are used in the place of 5 grammes of the Extract of Belladonna Leaves, in calculating the result of the volumetric test into terms of mydriatic alkaloids the product must be multiplied by 50 instead of 20. The quantity of alkaloids yielded from the 2 grammes of Extract employed should correspond to 2 p c

FLUIDEXTRACTUM SCOPOLA—The Fluid Extract of Scopola (*USP*) is prepared by exhausting dried Scopola Rhizome in No 40 powder with a mixture of 4 volumes of Alcohol (94.9 p c) and 1 volume of Water. It is required to contain 0.5 of a gramme of the mydriatic alkaloids from Scopola

Tests—The Fluid Extract is assayed by a process identical with that given by the *USP* for the assay of Fluid Extract of Belladonna Root, and the process is described under Extractum Belladonnæ Liquidum. A measured quantity (10 c c) of the Fluid Extract is employed, and it is required to contain an amount of mydriatic alkaloids corresponding to 0.5 p c w/v

Not Official

SCUTELLARIA.

The Herb of *Scutellaria lateriflora*, L., commonly known as Mad-dog Skull cap Official in U S

Scutellarin is a dry, light, greenish brown powder, not a pure, proximate principle. It may be prepared by precipitating a concentrated tincture with Water

Has been used in neuralgia, chorea, delirium tremens, and nervous exhaustion from fatigue or over excitement

Dose—1 to 5 grains = 0.065 to 0.32 gramme

A fluid extract (1 in 1) is also prepared, dose, $\frac{1}{2}$ to 1 fl drm = 1.8 to 3.6 c c U S has fluid extract 1 in 1

SENEGÆ RADIX.

SENEGAL ROOT

FR, POLYGALA DE VIRGINIE, GLR, SENEGAWURZEL, ITAL, POLIGALA VIRGINIANA, SIAM, POLIGALA DE VIRGINIA

The dried Root of *Polygala Senega*, L

Senega Root contains Saponin, and will therefore emulsify Oils, it also contains Methyl Salicylate

Medicinal Properties—A stimulating expectorant. Chiefly used in chronic bronchitis, especially if secretion be scanty, combined with Ammonium Carbonate, Ipecacuanha, and Squill

Official Preparations Infusum Senegæ, Liquor Senegæ Concentratus, and Tinctura Senegæ

Not Official—Fluidextractum Senegæ, Infusum Senegæ Concentratum, Syrupus Senegæ

Foreign Pharmacopœias—Official in all

Descriptive Notes—Typical Senega Root of good quality is yellowish-grey, curved irregularly, keeled on the inner or concave side, wrinkled longitudinally, and furnished with few branches. At the crown it is enlarged and shows traces of the bases of numerous slender stems. The fracture is short, and the horny translucent cortex is free from starch, and has an irregular one-sided development of liber tissue which forms the keel, the woody centre is white. The taste is sweetish at first, then acid, and the odour, especially in decoction, recalls that of Oil of Wintergreen (Methyl Salicylate). At intervals the root becomes scarce in commerce, and other varieties are offered, and the root is sometimes adulterated. A large root with a few principal branches at right angles, and with a crown sometimes 1 in (25 mm) across, with portions of the aerial stems attached, is known as Northern Senega, and is referred to the var. *latifolia*, T and G. Another variety, known as Southern Senega, resembles typical Senega, but has less cortex in proportion, and the keel is absent. It is referred to *Polygala Boykinii*, Nutt., or to *P. alba*, Nutt. Both of these differ from the official description in the absence of the keel. Occasionally other American medicinal roots have been found mixed with Senega either accidentally or purposely, but the only adulterant that might be confused with it is the rhizome of *Asclepias Vincetoxicum*, L., which closely resembles it in colour, and is apparently mixed with it in Europe. The distinct rhizomatous character of this adulterant at once distinguishes it, and the roots are crowded together and not contorted.

The distinctive features of the root under the microscope are the absence of starch, raphides, sclerenchymatous cells, and bast fibres, the presence of short tracheids, phloem parenchyma with oblique pores, collenchyma and drops of Oil in the parenchyma. The powdered root also readily gives a frothy solution when shaken with Water.

Tests—Senega Root yields from 2 to 5 p c of ash. The Swiss Pharmacopœia employs the detection of the presence of Methyl Salicylate in the ethereal Extract as a test of identity. An ash limit is not considered necessary.

Preparations

INFUSUM SENEGÆ INFUSION OF SENEGA

Senega Root, in No. 10 powder, 1, boiling Distilled Water, 20
Infuse half an hour, and strain (1 in 20)

Dose— $\frac{1}{2}$ to 1 fl oz = 1½ to 28 4 c c

Official in Belg. 3 of Fluid Extract in 100, Fl., Tisane de Polygalæ, 1 of Root in 100

LIQUOR SENEGÆ CONCENTRATUS CONCENTRATED SOLUTION OF SENEGA

A fluid extract (1 in 2) made with a mixture of 2 parts of Alcohol (20 p c) and 1 part of Alcohol (45 p c), by percolation (1 in 2)

Dose— $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

Tests—Concentrated Solution of Senega has a sp gr of 1 010 to 1 030, it contains from 10 to 20 p c w/v of total solids and from 18 to 22 p c w/v of Absolute Alcohol

TINCTURA SENEGÆ TINCTURE OF SENEGA

1 of Senega Root, percolated with Alcohol (60 p c), to yield 5 (1 in 5)

Dose— $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

Foreign Pharmacopœias—Official in Belg., 1 and 5, yielding 5 p c of dry residue, Mex., 1 in 5. Belg. has Polygalæ Extractum Fluidum, yielding 25 p c. dry residue, Dan. has Fluid Extract, 1 in 1

Tests—Tincture of Senega has a sp gr of 0 935 to 0 940, it contains from 3 to 6 p c w/v of total solids and about 55 p c w/v of Absolute Alcohol

Not Official

FLUIDEXTRACTUM SENEGÆ—100 of Senega Root in No. 40 powder, macerated and percolated with a mixture of 60 of Alcohol (95 p c), 30 of Water and 8 of Solution of Potassium Hydroxide, continue the percolation with a mixture of Alcohol (95 p c) 2, and Water 1, until drug is exhausted, reserve the first 80, evaporate remainder to a soft extract, which dissolve in reserved portion, and make up to 100 with more of the mixture of Alcohol (95 p c) and Water—*U S P*

This has been incorporated in the *B P C*

INFUSUM SENEGÆ CONCENTRATUM—Senega Root, in No. 20 powder, 40, Strong Solution of Ammonia, 0 5, Oil of Wintergreen, 0 15, Alcohol (90 p c), 1, Dilute Chloroform Water (1 in 1000), 3, *q s* to make 100. Mix the powder with the Strong Solution of Ammonia and sufficient menstruum to damp it evenly. Complete by repercolation. Dissolve the Oil of Wintergreen in the product. **Dose**— $\frac{1}{2}$ to 1 fl dm—*Fan*, and *Wright*, *P J* '06, 1 165 and '07, 1 622, *C D* '06, 1 252, and *Y B P* 1907, 251

This appears in the *B P C*

SYRUPUS SENEGÆ (U S)—Fluid Extract of Senega *U S* as above, 20, Syrup, 80

This has been incorporated in the *B P C*

Foreign Pharmacopœias—Official in Aust., Senega Root 5, Alcohol (90 p c) 5, Water *q s* to yield 45, Sugar 75, Belg., Fluid Extract 5, Syrup 95,

Dan, Senega Root 4, Water *q s* to yield 37, Sugar 68, Dutch, Ger, Jap and Russ, Senega Root 5, Alcohol (90 p c) 5, Water *q s* to yield 40, Sugar 60, Fr, Senega Root 1, Boiling Water 15, decant and add to each 10 of liquor 18 of Sugar, Hung, Senega Root 1, Dilute Alcohol 1, Water *q s*, to yield 10, Sugar 17, Ital, Senega Root 1, Water 12, Sugar 18, Mex, Extract 0 5, Alcohol (60 p c) 9 5, Syrup 90, Norw, Senega Root 4, Water *q s* to yield 10 Sugar 60 Span, Senega Root 15, Water *q s* to yield 360, Sugar 640, Swed Senega Root 5 Water *q s* to yield 37, Sugar 63, Swiss, a Fluid Extract is made by percolating 50 of the root with a mixture of Alcohol (90 p c) 1, and Water 5, addition of Ammonium Sulutum 5, evaporate to 50 and add 90 of Syrup

SENNA.

SENNA

FR, SENL, GER, SENNESBLATTFR, ITAL, SENA, SPAN, SEN DE ESPANA

The British Pharmacopoeia recognizes two kinds, **Alexandrian Senna**, the dried leaves of *Cassia acutifolia*, Delile, and **East Indian Senna**, the dried Leaflets of *Cassia angustifolia*, Vahl

When Senna is ordered in an official preparation either of the above may be used

U.S. has also an Indian and Alexandrian Senna, Ger an East Indian Senna. The different kinds of Senna, freed from stalks, are of nearly equal medicinal value

Medicinal Properties.—An efficient purgative in occasional or habitual constipation

Prescribing Notes.—As it produces griping and nausea, it is given with aromatics, such as Fennel in Compound Liquorice Powder, and Oil of Cloves in Syrup of Senna. The infusion is a suitable vehicle for Magnesium Sulphate and similar medicines

Dose.—10 to 30 grains = 0 65 to 2 grammes

Official Preparations.—Confectio Sennæ, Infusum Sennæ, Liquor Sennæ Concentratus, Syrupus Sennæ, and Tinctura Sennæ Composita. Contained in Pulvis Glycyrrhizæ Compositus. The infusion is used in the preparation of Mistura Sennæ Composita

Not Official.—Elixir Sennæ, Extractum Sennæ Leguminorum Liquidum, Extractum Sennæ Fructuum Fluidum, Infusum Sennæ Compositum, Lavement Purgatif and Acidum Catharticum

Foreign Pharmacopœias.—Official in all

Tests.—Senna contains from 8 to 14 p c of ash. Fourteen samples examined in the author's laboratory gave from 10 to 11 5 p c, with an average of 10 7 p c. The ash should be almost entirely soluble in Hydrochloric Acid

SENNA ALEXANDRINA. ALEXANDRIAN SENNA

The dried Leaflets of *Cassia acutifolia*, Delile

Descriptive Notes.—Alexandrian Senna arrives in this country in packages containing leaves roughly sorted into 1st and 2nd qualities, and the pods. The leaves are sifted and picked over on arrival in this country, in order to separate inferior leaves, broken twigs, and various impurities, and to grade the leaves into different sizes and

qualities The leaves, or more correctly leaflets, are opaque, of a light yellowish green tint, varying from $\frac{3}{4}$ to $1\frac{1}{4}$ in (19 to 30 mm) in length, lanceolate or oval, acute, mucronate, entire, unequal at the base, usually covered with a short fine pubescence, densest on the mid-rib, and the veins are conspicuous, especially on the under surface. The odour is tea-like but characteristic, and the taste mucilaginous, nauseous in a watery infusion. The characteristic feature is that the leaves are widest below the middle.

The leaves were formerly adulterated with Aigel leaves, but these are now rarely met with, these are readily distinguished by their equal base and minutely wrinkled surface. Alexandrian Senna leaves, as imported, vary chiefly in size, amount of debris and discoloured leaves present, the sifted and 'elect' Senna being worth about twice, and the hand picked leaves three to four times, the value of the crude drug as imported. The siftings, or small Senna, free from dust and sand, but containing about one-third of the weight of stalks and debris, are sold at about half the price of the crude drug. Occasionally the leaves of *Cassia obovata*, Collad, are found in Alexandrian Senna, and they are also occasionally imported separately, they are obovate in shape, and therefore easily recognised, but are considered to be less active than those of *C. acutifolia*. The pods of *C. acutifolia* are imported separately, the infusion is said to be milder in odour and flavour and slower in its action than that of the leaves, although equally effective. Powdered Senna is characterised by the 1-celled short hairs, slightly contracted below and tapering above, with thick and minutely papillose walls, by the cluster crystals in the parenchymatous cells, and the senate single prismatic crystals in the cells near the fibres of the veins, the polygonal epidermal cells and the long palisade cells.

Aigel leaves have 3-celled hairs, contain latex cells, and have short palisade cells.

SENNA INDICA. EAST INDIAN SENNA B P Syn — TINNEVELLY SENNA

The dried Leaflets of *Cassia angustifolia*, Vahl. From plants cultivated in Southern India.

Descriptive Notes — East Indian Senna leaflets are from 1 to 2 in (25 to 50 mm) in length, lanceolate and acute, with the greatest diameter near the middle, and rather less hairy than the Alexandrian kind. But the drug varies much in different samples, and like the Alexandrian needs sifting and sorting on arrival in this country. The different grades vary in size of the leaflets, freedom from discoloured leaflets and stalks and in their colour, the cultivated or Tinnevelly leaflets being usually greener, while those imported from Arabia, and known as Mecca Senna, are smaller, contain more stalks, are of a more faded or greyish-green tint, and often many discoloured leaflets are present.

The official description evidently indicates the better grades of the drug. Under the microscope the powder of the leaflets offers few distinguishing features from that of the Alexandrian kind.

According to Sayre (*PJ* (4) in p 458) the epidermal cells are smaller and more uniform in size and shape, with sharper angles, the cell of the Alexandrian being 40μ and those of East Indian Senna 35μ in diameter and the hairs are shorter and less curved and less numerous but the stomata are less round and more elongated or oval than the Alexandrian Senna, although they have also the appearance of two parallel cells near the ostiole, due to the two guard cells being below the epidermis. The pods of East Indian Senna are longer than those of Alexandrian Senna. Those of *C obovata*, Collad., differ in having short transverse ridges in the centre of the pod.

Preparations

CONFECTIO SENNÆ. CONFECTION OF SENNA *NO Syn* — LENTITIV ELECTUARY

Senna, 7, Comander Fruit, 3, Figs, 12; Tamarinds, 9, Cassia Pulp, 9, Plunes, 6, Extract of Liquorice, 1, Refined Sugar, 30. The Figs, Plunes, Tamarinds and Cassia Pulp are treated with Distilled Water and pulped through a sieve, when mixed with the other ingredients the yield should be 75, by weight (1 in 11 nearly)

Dose.—60 to 120 grains = 4 to 8 grammes.

Foreign Pharmacopœias—Official in all except Belg, Dan, Fr, Mex, Span and Swed, but differing in composition.

INFUSUM SENNÆ. INFUSION OF SENNA

Senna, 2 oz, Ginger, sliced, $\frac{1}{2}$ oz (55 grains), Distilled Water, boiling, 20 fl oz. Infuse 15 minutes, and strain (1 in 10)

From 20 fl oz of Infusion only about 14 fl oz drain out

Dose.— $\frac{1}{2}$ to 1 fl oz = 14.2 to 28.4 cc, as a draught, 2 fl oz = 56.8 cc

LIQUOR SENNÆ CONCENTRATUS. CONCENTRATED SOLUTION OF SENNA

20 of Senna, treated by continuous percolation with Distilled Water to yield 16 of fluid, which is heated to 180° F (82° C) for 5 minutes, and cooled. To this is added a mixture of Alcohol (90 p c) 2, and Tincture of Ginger 2 $\frac{1}{2}$. It should yield 20, by measure (1 in 1)

Dose.— $\frac{1}{2}$ to 1 fl dm = 1.8 to 3.6 cc.

Tests.—Concentrated Solution of Senna has a sp gr of 1.020 to 1.080, it contains from 12 to 18 p c w/v of total solids and about 18 p c w/v of Absolute Alcohol.

MISTURA SENNÆ COMPOSITA. COMPOUND MIXTURE OF SENNA *BP Syn*—BLACK DRAUGHT

Magnesium Sulphate, 5, Liquid Extract of Liquorice, 1, Compound Tincture of Cardamoms, 2, Aromatic Spirit of Ammonia, 1, Infusion of Senna, q s to yield 20 (1 of Magnesium Sulphate in 4)

BP 1885 contained Tincture of Senna, less Compound Tincture of Cardamoms, and no Aromatic Spirit of Ammonia.

Dose—As a draught, 1 to 2 fl. oz = 28 4 to 56 8 c c

Foreign Pharmacopœias—*Infusum Sennæ Compositum*—Dan, Coriander 2, Fructus Vitæ vinifere apocryna 5, Senna 10, Manna 25, Potassium Tartrate 8, boiling Water, *q s* to yield 720, Dutch, Senna 10, Anise 3, Water, *q s* to yield 80, Sodium Potassium Tartrate 10, Glycerin 10 Belg, Fluid Extract of Senna 10, Fluid Extract of Glycyrrhiza 5, Manna 20, Water 65, Ger and Jap, Senna 50, Boiling Water 450, Sodium Potassium Tartrate 50, Sodium Carbonate 1, Manna 100, Water, *q s* to yield 475, Alcohol (90 p c) 25, Norw, Senna 10, Coriander 2, Boiling Water, *q s* to yield 70, Manna 25, Potassium Tartrate 5, Russ, Senna 10, Boiling Water 60, Sodium Potassium Tartrate 10, Manna 15, Alcohol (90 p c) 3, Swed, Senna 10, Manna 20, Sodium Potassium Tartrate 10, Distilled Water, *q s* to yield 100, Swiss, Fennel 5, Senna 10, Manna 10 Sodium Tartrate 10, Water, *q s* to yield 100, U S, Senna 6, Manna 12, Magnesium Sulphate 12, Fennel 2, Boiling Water 80, Cold Water, *q s* to yield 100, Austr (Infusum Sennæ cum Manna) Senna 12, Water 100, Manna 15, Magnesium Carbonate 1, Fr (Apocryma Purgatif), Senna 2, Rhubarb 1, Sodium Sulphate 3, Manna 12, Boiling Distilled Water 20, Hung (Infusum laxativum), Senna 20, Boiling Water 160 Manna 30, Ital (Infusio di Senna con Manna) Senna 10, Water, *q s* to yield 150, Manna 25, Port (Infusio de Senna Composita) Senna 10, Anise 2, Manna 20, Sodium Potassium Tartrate 10, Boiling Water 100, Span (Infusion de Manna Laxante) Manna 60, Senna 20, Magnesium Sulphate 9, Water, *q s* to yield 300, also Infusion de Manna Purgante, Manna 90, Senna 12, Cinnamon Water 1, Water, *q s* to yield 300 Russ has also Infusum Sennæ Salinum, Senna 10, Boiling Water 100, Sodium Sulphate 10, Refined Honey 10

Tests—Compound Mixture of Senna has a sp gr of 1.115 to 1.120, it contains about 16 p c w/v of total solids and about 12 p c w/v of Absolute Alcohol

SYRUPUS SENNÆ SYRUP OF SENNA

50 oz of Sugar is dissolved with the aid of heat in 10 fl oz of a liquid extract of Senna (1 in 1), and when cool 10 minims of Oil of Coriander dissolved in 40 minims of Alcohol (90 p c) is added. It should yield 92 oz, by weight (1 in 1½)

Dose—½ to 2 fl drim = 1 8 to 7 1 c c

Foreign Pharmacopœias—*Syrupus Sennæ*—Dutch, Senna 10, Water, *q s* to yield 38, Sugar 62, Ger and Jap, Senna 10, Fennel 1, moisten them with Alcohol (90 p c) 5, pour on them Distilled Water 60, and extract in the cold for 12 hours, strain without pressing, boil the strained liquid, filter, after cooling dissolve in 35 of the filtrate 65 of Sugar, U S, Fluid Extract of Senna 250, Coriander Oil 5, Syrup, *q s* to make 1000 Syrupus Sennæ Compositus—Austr, Senna 10, Anise 1, Water 100, strain, and to each 10 add Manna 2, Sugar 15, Belg, Fluid Extract of Senna 75, Fluid Extract of Glycyrrhiza 15, Spirit of Anise 10, Syrup 900 Syrupus Sennæ Mannatus—Dan, Manna 150, Senna 100, Fennel 5, Ginger 5, Distilled Water, *q s* to yield 500, Sugar 500, Norw, Fennel 1, Ginger 1, Senna 10, Manna 15, Boiling Water, *q s* to yield 50, Sugar 50, Swed, Fennel 1, Senna 10, Manna 15, Distilled Water, *q s* to yield 50, Sugar 50 Hung, Syrupus Mannatus, Senna 30, Anise 3, Sodium Carbonate 3, Alcohol (70 p c) 30, Water, *q s* to yield 300, add Sugar 400, Manna 90 Ital, Sciroppo di Senna e Manna, Senna 15 Anise 2, Water, *q s* to yield 180, in which dissolve Manna 60, Sugar 200 Mex Jara bade San, Extract of Senna 2 5, Water 7 5, Syrup 90 Jap has also Syrupus Sennæ cum Manna, Senna 35, Fennel 2, pour on them boiling Water 850, set aside for 12 hours, express, in the expressed liquid 350, dissolve Manna 50, Sugar 400, allow to subside, decant the upper clear liquid, evaporate till it attains a syrupy consistence and strain

Tests.—Syrup of Senna has a sp gr of 1.290 to 1.320

TINCTURA SENNAE COMPOSITA. COMPOUND TINCTURE OF SENNA

Senna, 4 Raisins of commerce, freed from seeds, 2, Cassia Fruit, $\frac{1}{2}$, Coriander Fruit, $\frac{1}{2}$, Alcohol (45 p c), 20, by maceration (1 in 5)

Dose.— $\frac{1}{2}$ to 1 fl drm = 18 to 36 c c, for repeated administration, for a single administration, 2 to 4 fl drim = 71 to 142 c c

Foreign Preparations.—Official in Mex, 1 in 5 Fluid extract official in US 1 in 10 p c dry residue Belg and Dan have an alcoholic extract

Tests.—Compound Tincture of Senna has a sp gr of 0.985 to 0.995, it contains about 10 p c w/v of total solids and about 39 p c w/v of Absolute Alcohol

Not Official

ELIXIR SENNAE—16 fl oz of a liquor obtained from 16 oz of Senna by two macerations with a mixture of 4 of Alcohol (90 p c) and 12 of Distilled Water, is heated with 12 oz of Sugar to 200° F (93.3° C) for 10 minutes, 24 minims, Oil of Coriander, 2 minims, Tincture of Chloroform, 3 fl drim, and Alcohol (90 p c), 3 fl drim. Add them to the mixture, if necessary to 24 fl oz with Alcohol (60 p c).—*BPC Formulary* 1901, now incorporated in the *BPC* with the synonym Liquor Sennae Dulcis

Dose—1 to 3 fl drim = 36 to 106 c c

EXTRACTUM SENNAE LEGUMINORUM LIQUIDUM—20 fl oz of liquor obtained from 20 oz of Senna by two macerations with a mixture of 1 of Alcohol (90 p c) with 2 of Distilled Water. Heat to 200° F (93.3° C) for 10 minutes, and when cold add if necessary more of the mixture to make 20, filter.—*BPC Formulary* 1901, now incorporated in the *BPC* under the title **Extractum Sennae Liquidum**

Dose—1 fl drim = 36 c c

EXTRACTUM SENNAE FRUCTUUM FLUIDUM—Exhaust Senna Pods with cold Water and evaporate the resulting liquid *in vacuo*, so that 1 of Fluid Extract shall equal 1 of Senna Pods

INFUSUM SENNAE CONCENTRATUM—Senna Leaves broken small 80, Strong Tincture of Ginger, 5, Dilute Chloroform Water (1 in 1000) sufficient to make 100. Prepare by macero-expression. After completing the process, add the Strong Tincture of Ginger. Heat in a closed vessel by means of a water-bath to a temperature of 85° C, and maintain the heat for 5 minutes. **Dose**— $\frac{1}{2}$ to 1 fl drim, as a draught, 2 fl drim diluted with Water. *Farmer and Wright, PJ '06*, 165 and *OT*, 1622, *CD '06*, 1252 and *YBP* 1907, 211

The *BPC* includes a modification of this, the macero-expression is conducted with a mixture of Alcohol (90 p c), 1, Diluted Chloroform Water, 3, in place of the Chloroform Water as given above

INFUSUM SENNAE COMPOSITUM—Senna, 6, Manna, 12, Magnesium Sulphate, 12, Fennel, bruised, 2, Boiling Water, 80, Cold Water, q s to make 100. Upon the Senna, Manna and Fennel pour the boiling Water and macerate for 1 hour. Then add the Magnesium Sulphate in the infusion and pass cold Water through the strainer to make 100.—*USP*.

Dose—4 fl oz = 1136 c c

This has been incorporated in the *BPC*

LAVEMENT PURGATIF—Pour 500 of Water on to 15 of Senna Leaves and infuse half an hour, strain through a cloth, press, and dissolve in the fluid 15 of Sodium Sulphate.—*Fr*

ACIDUM CATHARTICUM—According to Stockman, Cathartic Acid is a coloured glucoside. In the free state it is easily decomposed. It acts locally as

an irritant and hence as a purgative when introduced into the alimentary canal
—*P J* (3) xv 751

Bougain and Bouchut, in a lengthy investigation on Cathartic Acid and Senna, conclude, 'As a general result of this inquiry it appears that the best preparation is the Infusion of Senna' —*P J* (3) ii 223

SERPENTARIÆ RHIZOMA.

SERPENTARY RHIZOME

The dried Rhizome and Roots of *Aristolochia Serpentina*, L., *Virginian Snakeroot*, or of *Aristolochia reticulata*, Nutt., *Texan* or *Red River Snakeroot*

From the southern parts of North America

Under the title of *Aristolochia* the dried Stem and Root of *Aristolochia indica*, L., are official in the *Ind* and *Col Add* for India and the Eastern Colonies

Medicinal Properties —A bitter stomachic See *Calumba*

Dose —10 to 15 grains = 0.65 to 1 gramme

Official Preparations —Infusum Serpentariæ, Liquor Serpentariæ Concentratus, and Tinctura Serpentariæ Used in the preparation of Tinctura Cinchonæ Composita

Not Official —Infusum Serpentariæ Concentratum

Foreign Pharmacopœias —Official in Mex., Port. and U.S.

Descriptive Notes —In the *BP* 1864 and 1867 the root of *Aristolochia Serpentina* was alone official, but the drug was rarely met with in commerce, the root of the Texan species, *A. reticulata*, being the article representing it in this country. In the *BP* 1885 and 1898 the latter species was made official, as well as the Virginian (*A. Serpentina*), and both are now obtainable. The latter has much more slender, matted, fibrous, furrowed roots, those of *A. reticulata* being longer, thicker, straighter and smoother. The rhizome of *A. Serpentina* is about 1 in (25 mm) long and $\frac{1}{2}$ in (3 mm) in diameter, bearing on its upper surface the remains of aerial stems, and the numerous slender, very interlacing roots are about 3 in (75 mm) long, yellowish brown in colour, have a bitter taste, and an odour recalling those of Camphor, Turpentine and Valerian. The rhizome of *Spigelia Marilandica*, L., resembles that of *A. Serpentina* in size and appearance, but it is not aromatic.

The characteristic microscopic features of Serpentary Root are the cuboid cells of the outer bark, the porous cuboid cells of the medullary rays, the oil cells in the mesophylleum, and the abundance of starch.

Tests.—Serpentary Rhizome contains from 7 to 10 p.c. of ash. An ash limit is not considered necessary.

Preparations

INFUSUM SERPENTARIÆ INFUSION OF SERPENTARY

Serpentary Rhizome, in No. 10 powder, 1, boiling Distilled Water, 20. Infuse 15 minutes, and strain (1 in 20)

Dose.— $\frac{1}{2}$ to 1 fl. oz. = 14.2 to 28.4 cc

LIQUOR SERPENTARIÆ CONCENTRATUS. CONCENTRATED SOLUTION OF SERPENTARY

A fluid extract (1 in 2) made with Alcohol (20 p c) (1 in 2)

Dose.— $\frac{1}{2}$ to 2 fl drms = 1.8 to 7.1 c c

Tests.—Concentrated Solution of Serpentry has a sp gr of 0.990 to 1.000, it contains about 5 p c w/v of total solids and about 18 p c w/v of Absolute Alcohol

A corresponding preparation, *Liquor Aristolochiæ Concentratus* (1 in 2), dose, 30 to 120 minims = 1.8 to 7.1 c c, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

TINCTURA SERPENTARIÆ. TINCTURE OF SERPENTARY

1 Serpentry Rhizome, percolated with Alcohol (70 p c), to yield 5 (1 in 5)

Dose. — $\frac{1}{2}$ to 1 fl drms = 1.8 to 3.6 c c

Foreign Pharmacopœias —Official in Mex and US, 1 in 5 US has also Fluid Extract

Tests.—Tincture of Serpentry has a sp gr of 0.895 to 0.900, it contains about 2 p c w/v of total solids and about 68 p c w/v of Absolute Alcohol

A corresponding preparation, *Tinctura Aristolochiæ* (1 in 5), dose, 80 to 60 minims = 1.8 to 3.6 c c, is official in the *Ind* and *Col Add* for India and the Eastern Colonies

Not Official

INFUSUM SERPENTARIÆ CONCENTRATUM

m No 20 powder, 10 Alcohol (90 p c), 25, Dilute Chl
qs to make 100 Prepare by
and '07, 1 622, CD '06 1 252, Farr and Wright, P J '06, 1 165
, 251
This appears in the B P C

SEVUM PRÆPARATUM.

PREPARED SUET

Fr, Suif de Mouton Purifié, GER, Hammeltalg, ITAL, Grasso di Montone, SPAN, Sebo de Carnero

A white, almost odorless, fatty substance, having a bland taste and unctuous to the touch. It is insoluble in Water. It is described in both *BP* and *USP* as the internal fat of the abdomen of the sheep, *Ovis Aries*, L., purified by melting and . . . The *USP* states that Prepared Suet should be kept in well-closed vessels impervious to fat. The *BP* gives no directions as to the precautions necessary in storage. The *USP* also states that it should not be used after it has become rancid.

Official Preparation —U . . . Unguentum Hydrargyri

Foreign Pharmacopœias —Official in all except Belg, Dutch and Russ, Fr (Suif de Mouton Purifié), Ital. (Grasso di Montone), Mex., Port and Span (Sebo)

Tests—Prepared Suet possesses a m p of from 45° to 50° C (113° to 122° F), the *BP* states from 44 4° to 48 9° C (112° to 120° F), the *USP* from 45° to 50° C (113° to 122° F), the *PG* from 47° to 50° C (116 6° to 122° F) The solidifying point is about 38° C (100 4° F), the *BP* states 37 8° C (100° F), the *USP* 37° to 40° C (98 6° to 104° F), the *PG* does not mention a solidifying point It is officially required to be freely soluble in Benzol, insoluble in cold Alcohol (90 p c), and slightly soluble in Ether or boiling Alcohol (90 p c) Acid, Saponification, and Iodine values might with advantage have been included The Acid value ranges from 0 8 to 2, the Saponification value from 192 to 195, the Iodine value from 33 to 46

It is officially stated that in India Benzotized Suet should be used in place of Benzotized Lard, see p 100

Not Official

SIMARUBA

BITTER SIMARUBA, OR MOUNTAIN DAMSON

The Root-bark of *Simaruba officinalis*, DC, from the West Indies

Medicinal Properties—A bitter tonic and astringent In large doses causes nausea Principally used in chronic forms of dysentery, may be combined with Opium

Dose—15 to 30 grains = 1 to 2 grammes

Foreign Pharmacopœias—Official in Dutch, Mex, Port and Swiss

SINAPIS.

MUSTARD

FR, MOUTARDI, GER, SINISAMEN, ITAL, SPINAPI NFRA, SPAN, MOSTAZA

The powdered and mixed dried ripe Seeds of *Brassica nigra* and *Brassica alba*

The whole virtue of Mustard depends upon the fact that when mixed with Water, Allyl Thiocarbimide (Volatile Oil of Mustard) is formed This compound is produced by the action of Myrosin upon Myronic Acid in the same way in which the Emulsin and Amygdalin react in the formation of Volatile Oil of Bitter Almonds Black Mustard contains Myrosin and a large excess of Myronic Acid, and so is in itself able to produce the Volatile Oil to some extent White Mustard contains Myrosin but no Myronic Acid, and so can by itself produce none of the Volatile Oil The best result is obtained by mixing the black and white variety in such proportions that the Myrosin and the Myronic Acid will balance each other

Medicinal Properties—A powerful stimulant and sialagogue The powder is taken internally as a condiment, a tablespoonful in a tumblerful of warm Water acts as a prompt emetic, used externally in form of poultice or chaeta, as a rubefacient and counter-irritant in pneumonia, pleurisy, muscular rheumatism, pericarditis, bronchitis, colic, gastralgia, vomiting and neuralgia, as a sitz-bath in amenorrhœa.

Official Preparations—Charta Sinapis, Linimentum Sinapis, and Oleum Sinapis Volatile

Not Official—Applicatio Sinapis, Cataplasma Sinapis, Unguinum Sinapis, Thiosinamin and Fibrolvsin

Foreign Pharmacopœias—Official in Part (Mostaïde)

Tests—Mustard yields about 5 p c of ash. It is officially required that a cold decoction should not be rendered brown by Boric Acid Solution, indicating the absence of Turmeric, nor should it yield a distinctive reaction with the tests for Starch. The test for Turmeric may be conveniently carried out with Methylated Spirit as a solvent instead of Water. About 1 gramme of the specimen should be boiled with Methylated Spirit, filtered, the filtered liquid concentrated and tested with Boric Acid Solution. The Iodine test for the presence of Starch is rendered negative, owing to the ready absorption of the Iodine by the volatile Oil developed on the addition of Water. In contrast to the Iodine test for Starch on the powdered seeds of the White or Black Mustard, the *U S P* directs that 1 gramme of the powdered Mustard should be exhausted by slow percolation with Alcohol (94.9 p c), the marc mixed with 200 cc of Water and heated to boiling, adding after cooling sufficient cold Water to make the mixture measure 1000 cc, the addition of 4 cc of Tenth-normal Volumetric Iodine Solution should not produce a dark blue colour. The *German Pharmacopœia* includes a test for the determination of the ethereal Oil. A quantity of 5 grammes of the powdered Mustard is digested in 100 cc of Water at a temperature of 20° to 25° C (68° to 77° F). The stoppered flask is allowed to remain at rest for 2 hours, the contents are then added 20 cc of Alcohol (90 p c) and 2 cc of Olive Oil, the flask connected with a well-cooled condenser, and the mixture is distilled. The first 40 to 50 cc of distillate is collected in a graduated flask of 100 cc capacity, containing 10 cc of Ammonia Solution, and 20 cc of Tenth-normal Silver Nitrate Solution are added. The mixture is then diluted with Water to the mark on the neck and allowed to stand in a stoppered flask for 24 hours, with intervals of frequent shaking. To a measured quantity (50 cc) of the clear filtrate is then added 6 cc of Nitric Acid and 1 cc of Ferric Ammonium Sulphate Solution, and the mixture is titrated with Tenth-normal Volumetric Ammonium Rhodanate Solution until a red coloration appears, not more than 7.2 cc of this solution should be required, each cc of Tenth-normal Volumetric Silver Nitrate Solution absorbed represents 0.0049575 gramme of Allyl Thiocarbimide. The amount of Water present should not exceed 5 p c.

SINAPIS ALBÆ SEMINA. WHITE MUSTARD SEED

The dried ripe Seeds of *Brassica alba*

Descriptive Notes—The White Mustard Seed of commerce is often a mixed article, depending upon its geographical source, but the official article is the seed of *Brassica alba*, Boiss. (*Sinapis alba*, L.) The seed is yellowish, about $\frac{1}{12}$ in (2 mm) in diameter and $\frac{1}{10}$ of a grain in weight (*Pharmacographia*, p 69). The testa appears to be

smooth, but under a good lens is seen to be minutely and reticulately pitted. The embryo is of a bright, pure yellow colour and oily. It has no pungent odour until triturated with Water. The outer layer of epidermal cells contains mucilage and swells up rapidly in Water, a property which is sometimes taken advantage of for drying bottles intended to contain oily liquids, and from which it is necessary to remove the last trace of moisture. This is quickly effected by shaking a small quantity of White Mustard Seed in the bottle. The seeds of a false White Mustard, named by Huiz *Brassica albifolia*, have been substituted for those of *B. alba*, but they are slightly more ochreous in colour, the hilum is darker than the rest of the seed, and the epidermal cells are not mucilaginous.

The pods containing White Mustard Seed are spreading, and have busily hairs, and half their length is occupied by a flat, voiny beak. In Black Mustard the pods are erect and glabrous, and the short beak is slender and quadrangular.

The seeds of *S. glauca*, Roxb (*B. campestris*, L.), resemble those of *B. alba* and probably form part of the White Mustard Seed of India.

SINAPIS NIGRÆ SEMINA BLACK MUSTARD SEED

The dried ripe Seeds of *Brassica nigra*, Koch

Descriptive Notes - The Black Mustard Seed of commerce is usually brown rather than black, and often consists chiefly of the slightly larger and more oblong Indian or Sarepta Mustard Seeds, which are the produce of *Brassica juncea*, Cass. The seeds of *Brassica nigra*, Koch, which are official under the name of Sinapis Nigræ Semina, are $\frac{1}{16}$ in (1 mm) in diameter and $\frac{1}{10}$ of a grain in weight and of a dark reddish-brown or greyish-brown colour, and have at first a bitter and quickly afterwards a very pungent taste. Although reticulated with minute pits when dry, the surface of the seed appears smooth when wetted, owing to the development of mucilage from the outer walls of the epidermal cells. The whitish pellicle, which gives the greyish tint to some of the seeds, is attributed to rain during the ripening, and depreciates the value of the seed in the market; this pellicle is formed from hexagonal tabular cells. The seed contains about 20 per cent of fixed Oil, which is expressed and sold separately as a remedy for rheumatism, although it is used in Russia like the best Olive Oil (*Pharmacographia*). The ferment Myrosin is coagulated at 140° F (60° C), so that boiling Water must not be used in making Mustard plasters. The prepared Mustard leaves must be kept quite dry, or the ferment gradually acts in the presence of the moisture attracted from the air, and the Mustard leaves lose their efficacy. Powdered Mustard that has become damp is liable to become attacked by the cheese mite *Tyroglyphus Siro*, Geiv. The distinctive microscopic features of powdered Mustard are the absence of Starch and raphides, the mucilaginous epidermal polygonal cells, appearing striated when wet, large collenchymatous cells, and the yellow sclerenchyma of *S. alba*, and the dark yellowish-brown sclerenchymatous cells of *S. nigra*, some of which are rather longer and

form a hexagonal network, and the small, irregular aleurone grains (0.017 mm long and 0.008 mm broad) containing minute globoids and drops of fixed oil

Preparations

CHARTA SINAPIS. **MUSTARD PAPER**

Extract by Benzol the fixed Oil from a mixture of equal weights of bruised Black and White Mustard Seeds, dry, and reduce to No. 60 powder; mix 75 grains of it with 5 fl. drim. of Solution of Indiarubber, and spread by means of a suitable brush over about 30 sq. in. of one side of a piece of cartridge paper. Allow it to dry by exposure to the air.

Foreign Pharmacopœias.—Official in Austri, Belg, Dutch, Norw., Fr. (Sinapismes en Feuilles), Dan., Ger., Hung. and Swed. (Charta Sinapisata), Ital. (Carta Senipata), Mex. (Sinapismes de Papel), Spai. (Papel Sinapico), Russ. (Charta Sinapina), Swiss and U.S.

LINIMENTUM SINAPIS. **LINIMENT OF MUSTARD**

Volatile Oil of Mustard 1½ fl. drim., Camphor, 120 grains, Castor Oil, 5 fl. drim. Alcohol (90 p.c.), 4 fl. oz.

Now about 1 in 27 instead of 1 in 40, and Ethereal Extract of Mezeleum omitted.

As the volatile Oil quickly disappears on keeping it is better to keep the other ingredients ready mixed, and to add the Mustard Oil when required.

Spiritus Sinapis.—Austri and Hung. Oil 1 Spirit 50, Belg, Ger., Jap., Russ., Swed. and Swiss, Oil 1, Spirit 49, Mex. (Linimento de Mostaza Compuesto) 1 in 38. All by weight.

OLEUM SINAPIS VOLATILE. **VOLATILE OIL OF MUSTARD**

An almost colourless, or pale yellow, highly refractive, limpid, oily liquid, possessing a very characteristic, penetrating, pungent, and excessively irritating odour. It is the volatile oil distilled from Black Mustard Seeds after maceration with Water.

The *B.P.* describes the Oil as distilled from Black Mustard Seeds after maceration with Water. The *U.S.P.* describes it as a volatile Oil obtained from Black Mustard (freed from its fatty oil by maceration with Water and subsequent distillation). The *P.G.* describes it as a volatile Oil obtained by distillation from powdered Mustard Seeds which have been macerated in Water. The *B.P.* do not require it to contain any definite percentage of Allyl C_6H_5 . The *U.S.P.* requires that it shall contain not less than 92.4 to 99 p.c. The *P.G.* requires that it shall contain from 92.4 to 99 p.c.

It should be kept in well-stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from contact with the light. Its vapour is intensely irritating, and the greatest caution should be exercised in handling the oil.

Solubility.—1 in 50 of Water, readily in Alcohol (90 p.c.) and Ether.

Medicinal Properties.—Applied to the skin, it produces almost instant vesication, but when diluted it forms a useful counter-irritant application.

Foreign Pharmacopœias.—Official in all except Dan.

Tests—Volatile Oil of Mustard has a sp gr of 1.015 to 1.030. The *BP* states 1.018 to 1.030, the *USP* 1.013 to 1.020 at 25° C (77° F), the *PG* 1.018 to 1.025. It boils at 145° C (298.4° F), and distils between this temperature and 153° C (307.4° F). It is officially stated to distil between 147.2° and 152.2° C (297° and 306° F), the *USP* states that if a portion of the Oil be heated in a flask connected with a well cooled condenser it should distil completely between 145° and 152° C (298.4° and 305.6° F). The *PG* states that the boiling point is between 148° and 152° C (298.4° and 305.6° F); with the simple statement of these physical characteristics the *BP* is content. The *USP* and the *PG* state that if to 3 grammes of the Oil 6 grammes of Sulphuric Acid be gradually added, keeping the liquid cool, the mixture upon subsequent digestion will evolve Sulphur Dioxide gas, that it will remain of a light yellow colour, and although at first clear will afterwards become thick and occasionally crystalline, and will lose its pungent odour. These two Pharmacopœia also mention that when diluted with 5 times its volume of Alcohol (94.9 p.c. *USP*, 90 p.c. *PG*) the addition of a drop of Ferric Chloride Test Solution shall produce no blue or violet coloration. The *BP* does not introduce a method of assay. The *USP* requires that it shall contain not less than 92 p.c. of Allyl Iso-thiocyanate as determined by a process of which the following are the essential details—A quantity of about 2 grammes of the Oil is accurately weighed and diluted with sufficient Alcohol (94.9 p.c.) to produce a solution, 50 c.c. of which shall represent 1 gramme of the Oil. A measured quantity of 5 c.c. of this solution is transferred to 100 c.c. measuring flask, and 30 c.c. of Tenth-normal Volumetric Silver Nitrate Solution and 5 c.c. of Ammonia Solution added. After well stoppering the flask the mixture is set aside in a dark place for 24 hours, then heated in a water-bath at a temperature of 80° C (176° F) for half an hour, with frequent intervals of shaking, the contents of the flask are diluted to the 100 c.c. mark and filtered. A measured quantity of 50 c.c. of the filtrate is mixed with 4 c.c. of Nitric Acid and a few drops of Ferric Ammonium Sulphate TS and sufficient Tenth-normal Volumetric Potassium Sulphocyanate Solution to produce a permanent red colour is added, not more than 5.6 c.c. of such solution should be required. Each c.c. of Tenth-normal Volumetric Silver Nitrate Solution absorbed corresponds to 0.00492 gramme of Allyl Iso-thiocyanate. The *PG* process is essentially as follows—A measured quantity of 5 c.c. of a 1 in 50 w/v solution of the Oil in Alcohol (90 p.c.) is transferred to a 100 c.c. stoppered measuring flask, 50 c.c. of Tenth-normal Volumetric Silver Nitrate Solution and 10 c.c. of Ammonia Solution added, and the flask is well stoppered and the mixture allowed to stand for 24 hours, with intervals of vigorous shaking. It is then diluted with Water to the 100 c.c. mark. A measured quantity of 50 c.c. of the clear filtrate is mixed with 6 c.c. of Nitric Acid and 1 c.c. of Ferric Ammonium Sulphate Solution, and sufficient Tenth-normal Volumetric Ammonium Rhodanate Solution added to produce a red coloration, from 16.6 to 17.2 c.c. shall be required, *PG*.

Owing to the results yielded by the *P G* method being invariably too low, the following modifications of the *P G* method are suggested by Schimmel in the *Berichte* for April 1908 —A measured quantity of 1 or 5 cc of a 1-50 solution of the Oil in Alcohol (90 pc) is placed in a 100 cc stoppered measuring flask with 10 cc of Ammonia Solution and 50 cc of Tenth-normal Volumetric Silver Nitrate Solution, and after closing the flask with a cork provided with a reflux tube 1 metre long it is heated for about 1 hour on a water-bath which is kept at about 100° C. boiling. After cooling to 15° C. and diluting with Distilled Water to the mark, it is filtered, a measured quantity of 50 cc of the clear filtrate, after addition of sufficient Nitric Acid to produce a faintly acid reaction, is titrated with Tenth-normal Volumetric Ammonium Rhodanate Solution until the appearance of a red colour, 1 cc of Ferric Ammonium Sulphate Solution being added as an indicator, from 16.6 to 17.2 cc of the solution should be necessary. The *P G* figures require that it shall contain from 92.4 to 99.0 pc, calculated as Allyl Iso-thiocyanate. It is preferably determined by the above process.

The more generally occurring impurities are Ethyl Alcohol, Petroleum, Chloroform, fatty oils, Carbon Bisulphide, Phenols. The requirement that the Oil shall distil between 147.2° and 152.2° C (297° and 306° F), and that the first and last portions of the distillate should have the same sp gr as the original Oil, precludes the presence of Ethyl Alcohol, Petroleum, Chloroform, fatty oils and Carbon Bisulphide. For the actual determination of Carbon Bisulphide, Schimmel recommends the following process —A weighed quantity of 20 to 25 grammes of the Oil are heated on a water-bath while a slow current of air is passed through the Oil, the vapour of Carbon Bisulphide which is thus carried off is cooled by passing through a condenser and conducted into an alcoholic Potassium Hydroxide Solution, where it is converted into Potassium Ethyl Xanthate. After the neutralisation of the alkali solution, sufficient Tenth-normal Volumetric Copper Sulphate Solution is added until a drop produces a reddish-brown colour with Potassium Ferrocyanide. From the amount of Copper Solution consumed the percentage of Carbon Bisulphide present can be ascertained. 1 cc of Tenth-normal Volumetric Copper Sulphate Solution corresponds to 0.0152 gramme of Carbon Bisulphide. The volumetric process may be supplemented by a gravimetric one, the precipitate of Cuprous Ethyl Xanthate may be collected on a filter, washed, dried and heated to a red heat in a crucible and the residue of Cupric Oxide weighed. 1 gramme of the Oxide corresponds to 1.198 grammes of Carbon Bisulphide. Phenols, if present, may be detected by the test with Ferric Chloride T.S. described above.

Not Official

APPLICATIO SINAPIS —Volatile Oil of Mustard, 4 minims, Eau de Cologne, 1 fl oz. Mix.

A good application in acute catarrh of the middle ear, to be applied behind the ear on the side of a brush or absorbent Wool.

CATAPLASMA SINAPIS —Mustard, in powder, 2½ oz or 95, Linseed Meal, 2½ oz, boiling Water and Water, of each a sufficiency —*B P* '85

Mix the Mustard with 2 or 3 oz of lukewarm Water, mix the Linseed Meal with 6 to 8 oz of boiling Water, add the former to the latter and stir together.

Crushed Linseed, 28, Mustard, 2, Water, *q s* to make 100 — *B P C*

This is similar to the following —

Make a Linseed Poultice by adding 4 parts of crushed Linseed to 10 parts of boiling Water, and for every 4 oz of crushed Linseed employed add $\frac{1}{2}$ oz of Mustard, previously rubbed to a smooth paste with a little cold or tepid Water — *St Thomas*

INFUSUM SINAPIS — Mustard, 2 dm, boiling Water, 1 fl oz, strain. It relieves obstinate hiccuph.

THIOSINAMIN (Allyl thio carbamide) — White, glistening crystals, generally odourless, but sometimes possessing a faint, garlic odour, soluble 1 in 17 of Water, 1 in 2 of Alcohol (90 p c) and soluble in Ether. It has been found useful for softening scar tissues and the removal of fibrous structure of the oesophagus, etc., and has also been used in the treatment of lupus.

Employed in the form of a 15 to 20 p c alcoholic solution, $\frac{1}{2}$ to 1 syringeful being injected between the scapule or as an 8 p c solution in Water containing 20 p c Glycerin, 20 minims being injected, in divided portions, in the neighbourhood of the growth — *B M J E* '02, 1 91, '04, 1 75, *B M J* '03, 1 656, *L* '08, 1 785, *C D* '02, 1 538, *P J* '02, 1 201.

Fibrolysin — In arthritis deformans with contractures, and in old-standing urethral stricture — *Pr* '07, 1 427. Successful in treating gastric adhesions — *F T* '07, 88.

In perigastric adhesions, pyloric structure, hourglass stomach, and conditions in which there is new formation of connective tissue, it may be given (*B M J* '05, 1 811) hypodermically in the form of a 10 p c solution with 70 parts of Water and 20 parts of Alcohol. It has been successfully used (*B M J* '06, 1 379) in the treatment of hypertrophy of the pylorus. 10 minims of a 10 p c solution in Alcohol injected daily for 1 week, then every other day for 14 days, then 3 times a week for 6 weeks, then twice a week for a month, and then 15 minims once a week for another 3 months.

SODIUM.

SODIUM

Na, eq 22 88

A light, soft metal, exhibiting a silvery metallic lustre when freshly cut, but which rapidly oxidises in contact with air. It should be preserved under mineral Naphtha in well-stoppered glass bottles.

Metallic Sodium is not official in either the *U S P* or the *P G*.

The only direct official preparation of Sodium is **Liquor Sodii Ethylatis** *See* **SODII ETHYLATIS LIQUOR**, p 1115.

Tests — Sodium has a sp gr of 0 97. It possesses a strong affinity for Oxygen and rapidly oxidises in the air when cut. When thrown on to cold Water it instantly fuses to a globule without combustion and traverses the surface in all directions, when thrown on to hot Water, however, or if its movements be circumscribed, combustion of the evolved Hydrogen ensues. The Water acquires an alkaline reaction towards red Litmus paper. It is officially mentioned that Water and Alcohol (90 p c) are vigorously attacked by it, Hydrogen being simultaneously evolved, the metal is almost entirely dissolved, leaving little or no insoluble residue. It is required to indicate at least 97 46 p c of metallic Sodium as determined by very

cautiously adding 1 gramme of Water, and titrating the resulting solution with Normal Volumetric Sulphuric Acid Solution, at least 42.6 cc are officially required for one cc of soda. The *BP* does not mention which indicator of neutrality should be used, but Phenolphthalein Solution may be most conveniently employed. When evaporated to dryness, the residue yields a brilliant yellow coloration when moistened with Hydrochloric Acid and introduced on a platinum wire into a non-luminous flame.

Not Official SODA CAUSTICA.

White, hygroscopic pencils, or sticks, possessing a crystalline structure, or as a white, crystalline, or in powder, or in fused masses. It should be kept in well-closed hard glass bottles and exposed as little as possible to the air, as it is liable to absorb moisture and Carbonic Anhydride from the air. It is necessary to exercise very great caution, as it has a strong irritating effect from the powder is very irritating to the eyes and nose. The *BP* does not state any requisite percentage of pure Sodium Hydroxide, the *USP* requires that it shall contain not less than 90 p. Hydroxide, and not more than 2 p. of other exception of Water. It is not official in the *PG*. A pure Sodium Hydroxide is also official in the *BP*. Appendix, the latter is only required in testing for traces of Aluminium. It is officially directed to be obtained by dissolving Sodium Hydroxide in water, and the solution evaporating to dryness in a silver dish, as it occasionally contains occasional evaporation.

Foreign Pharmacopœias—Official in Austr (Natrium hydroxydatum), Din, Dutch and Swed (Hydras Natricus), Ital (Soda Caustica), Jap (Natrium Causticum), Port (Hydrato de Soda), Span (Hidrato Sodico), Swiss (Natrium Hydricum), US (Sodium Hydroxidum).

The Solution is official in Austr (Natrium Hydroxydatum Solutum) (15 p.), sp gr 1.169 to 1.172, Dutch (Solutio Hydriatis Natrii) (13½ p.), sp gr 1.155, Fl (Souda Caustique Liquide) (about 30 p.), sp gr 1.392, Ger (Liquor Natrii Caustici) (15 p.), sp gr 1.168 to 1.172, Hung (Natrium Hydroxydatum Solutum) (32 p.), sp gr 1.185, Port (Hydrato de Soda Liquido), sp gr 1.33, Span (Solucion de Sosa Caustica) (30 p.), sp gr 1.33, Swed (Solutio Hydriatis Natrii) (25 p.), sp gr 1.275 to 1.285, Swiss (Natrium Hydricum Solutum) (30 p.), sp gr 1.33, US (Liquor Sodium Hydroxidi) (about 5 p.), sp gr 1.056 at 25°C (77°F).

Antidotes—Same as Liquor Potassæ, p. 930.

Pasta Londinensis—Caustic Soda, Unslaked Lime, equal parts, reduced to a fine powder, and kept in a well-closed bottle. To be made into a paste with Water when required.

Tests—Sodium Hydroxide dissolves readily and completely in Water, the solution even when very highly diluted has a strong alkaline reaction towards red Litmus paper. Sodium Hydroxide may be readily determined by direct titration with Normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. The quantity of Carbonate present in commercial samples can be determined by the method when working on a large scale, or by the method of the *USP*. When Merck's Orange Solution is employed as an indicator, the Carbonate is also estimated. 1 cc. of Normal Volumetric Sulphuric Acid is equivalent to 0.03976 gramme of pure anhydrous Sodium Hydroxide. The *USP* requires that it shall contain not less

than 90 p.c. of pure anhydrous Sodium Hydroxide, as determined by introducing about 1 gramme of the salt into a stoppered weighing bottle and accurately ascertaining its weight, dissolving in about 50 c.c. of Water, and titrating the Solution with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as an indicator of neutrality, the number of c.c. of Normal Volumetric Sulphuric Acid Solution required multiplied by 3.976, the product divided by the weight of Sodium Hydroxide taken, the quotient represents the percentage of pure anhydrous Sodium Hydroxide present. When neutralised with Hydrochloric Acid, the product when introduced on a platinum wire into the non luminous flame affords a brilliant yellow coloration. The *U.S.P.* states that when heated to about 525° C (977° F) it melts to a clear oily liquid and is slowly volatilised unchanged at a bright red heat.

The more generally occurring impurities are organic matter and insoluble impurities, heavy metals such as Arsenic, Copper, Lead, Iron and Zinc, Potassium Carbonate, Silicate, Chlorides and Sulphates. The 1 in 20 aqueous solution should be perfectly clear and colourless, indicating the absence of organic matter and insoluble impurities. When acidulated with Hydrochloric Acid it should yield no coloration or precipitate with Hydrogen Sulphide, nor should any coloration or precipitate ensue upon the subsequent addition of Ammonia Solution, indicating the absence of heavy metals. A 5 p.c. aqueous solution after acidification with Acetic Acid should yield no precipitate on the addition of Tartaric Acid, indicating the absence of Potassium. It should not yield more than a faint effervescence when a slight excess of diluted Sulphuric Acid is added to 10 c.c. of a 10 p.c. solution, indicating the limit of Carbonate. When 0.7 of a gramme of Sodium Hydroxide is dissolved in 1.5 c.c. of Water, this solution should not yield more than a slight white precipitate, within 10 minutes, when added to 10 c.c. of Alcohol (94.9 p.c.) indicating a limit of Silicate. A 5 p.c. solution when acidified with Nitric Acid should not yield more than a faint turbidity with Silver Nitrate or Bismuth Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. The *B.P.* includes also Aluminium and Phosphates as likely impurities. Aluminium, if present, may be detected by neutralising the Hydroxide with Hydrochloric Acid, adding Ammonium Chloride and Ammonia Solution and boiling, no white flocculent precipitate should be produced. A 5 p.c. aqueous solution should not afford a yellow precipitate when acidified with Nitric Acid and warmed with Ammonium Molybdate T.S., indicating the absence of Phosphates. The purified Sodium Hydroxide of the *B.P.* is required to yield no characteristic reactions with the tests for Phosphates or Sulphates, and not more than the slightest reactions with the tests for Carbonates, but need not necessarily be quite free from Aluminium.

LIQUOR SODII HYDROXIDI—Solution of Sodium Hydroxide. Purified Sodium Hydroxide, 200 grammes, Distilled Water, sufficient to produce 1000 c.c. The purified Sodium Hydroxide is dissolved in a portion of the Distilled Water, the solution made up to 1000 c.c. and filtered.

Tests—Solution of Caustic Soda *B.P.* has a sp. gr. of about 1.175. It contains about 18.0 p.c. of pure anhydrous Sodium Hydroxide, as determined by titrating a measured quantity of the Liquor with Normal Volumetric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. The *B.P.* does not mention any requisite percentage, the *U.S.P.* Liquor has a sp. gr. of about 1.056 at 25° C (77° F) and is required to contain about 5 p.c. of pure anhydrous Sodium Hydroxide. About 25 c.c. of Normal Volumetric Sulphuric Acid Solution are stated to be necessary to neutralise 20 c.c. (19.9 grammes) of the solution, using Methyl Orange T.S. as an indicator of neutrality. 1 c.c. of Normal Volumetric Sulphuric Acid indicates 0.2 p.c. of absolute Sodium Hydroxide. The *P.G.* Liquor has a sp. gr. of 1.168 to 1.172, it is required to contain about 15 p.c. w/w of absolute Sodium Hydroxide. The *B.P.* solution naturally should be free from such impurities as are precluded from Sodium Hydroxide or purified Sodium Hydroxide. The *U.S.P.* Liquor is required to answer the same reactions and tests as an aqueous solution of Sodium Hydroxide. The *P.G.* Liquor is required to be free from Carbonates, to contain only traces of Chlorides and Sulphates, to be free from Nitrates, and to contain only traces of Aluminium.

SODA TARTARATA.

SODIUM POTASSIUM TARTR

D P Syn—TARTARATED SODA, TARTRATE OF POTASSIUM
U S Syn—TARTARUS NATRONATUS

ALLG
 ITF.



FR, TARTRATE DROIT DE SODIUM
 TARTRAT, ITAL, TARTRATO
 POTASSICO

ALUMNAPRIUM-
 TARTRATO SODICO-

Colourless, transparent crystals, or a white, odourless powder, having a slightly bitter taste. Prepared by neutralising the acid radicle with Sodium Carbonate and recrystallising.

It is soluble in Water, and is used in vessels and exposed as little as possible to the air, which has a slight tendency to effloresce.

It is soluble in Water, and is used in vessels and exposed as little as possible to the air, which has a slight tendency to effloresce.

Medicinal Properties.—A mild purgative, well suited for constipation associated with gout and hepatic dyspepsia. It is not aperient in small doses, its action then being diuretic, antilithic, and to render the urine alkaline.

Dose—120 to 240 grains = 8 to 16 grammes

Official Preparation—Pulvis Sodæ Tartarata Effervescens

Foreign Pharmacopœias—Official in Aust and Hung (Kalium Natrio-tartaricum), Belg (Kalium Natrium Tartaricum), Dan, Norw and Swed (Tartarus Natrio-kalicus), Dutch (Tartarus Kalico-natricus), Fr (Tartate Droit de Potassium et de Sodium), Ger and Swiss (Tartarus Nationatus), Ital (Tarttrato Sodico-Potassico), Jap and Russ (Natrio-Kalium Tartaricum), Mex (Tarttrato de Potasio y Sodio), Port (Tarttrato de Potassa de Soda), Span (Tarttrato Sodico-Potassico), US (Potassii et Sodii Tartras)

Tests.—Tartarated Soda when heated fuses to a more or less colourless liquid and loses its Water of crystallisation, equivalent to 25.5 p.c. At a higher temperature it gradually becomes brown, and when still more strongly heated evolves an odour of burnt Sugar and leaves a black residue possessing a strong alkaline reaction. It dissolves readily in Water, forming a colourless solution possessing a faintly alkaline reaction towards Litmus paper. The *P G* states that it is neutral towards Litmus paper. The *U S P* states that the aqueous solution does not effect Phenolphthalein Solution. The *B P* makes no mention of its reaction towards any indicator of neutrality. When incinerated and the residue is dissolved in diluted Hydrochloric Acid it yields a solution which answers the distinctive tests of Potassium and Sodium given under those headings. The aqueous solution affords, with Calcium Chloride Solution, a white granular precipitate soluble in a cold, moderately concentrated Potassium Hydroxide Solution, being again reprecipitated on boiling, with Silver Nitrate Solution it yields a white precipitate soluble in Nitric Acid and in

Ammonia Solution, and if just sufficient Ammonia Solution be added the precipitate redissolves and the mixture yields on boiling in a perfectly clean test tube a mirror of metallic Silver. When a moderately concentrated solution of the salt is acidulated with Acetic Acid and mixed with a concentrated Potassium Acetate Solution it affords, when well stirred, a white precipitate, the precipitation being more pronounced on the addition of Alcohol (90 p c). When acidified with Acetic Acid it yields, on the addition of a drop of Ferrous Sulphate Solution, a few drops of Hydrogen Peroxide Solution and an excess of Potassium Hydroxide Solution, a purple or violet coloration. It is officially required to contain 98.0 p c of pure crystallised Sodium Potassium Tartrate, as volumetrically determined by the method given below under Volumetric Determination. The *U.S.P.* requires the salt to contain not less than 99 p c of pure Potassium and Sodium Tartrate, the process of determination being also a volumetric one and appearing below. The *P.G.* does not include a method of determination, nor does it state the amount of pure salt which it is requisite for a specimen to contain. As regards impurities, the *B.P.* does not mention any substances as likely impurities, the more generally occurring are Lead, Copper, Iron, Ammonium salts, Calcium, Sulphates and Chlorides. Lead, Copper and Iron may be detected by the Hydrogen Sulphide test, Ammonium salts by the Potassium or Sodium Hydroxide test, Calcium by the Ammonium Oxalate test, Sulphates and Chlorides by the Barium Nitrate and Silver Nitrate test described below. The absence of tests for impurities has apparently escaped the notice of those responsible for the *Report of the Committee of Reference in Pharmacy*, as no recommendation for their inclusion appears, and a limit of Lead as an impurity should have been included, not only in the present instance, but in the case of all Tartrates and Citrates. Standards have been suggested (*C.D.* '08, 1796) of 10 parts per million for Lead, and 2 parts per million for Arsenic. A suitable limit for Lead in Tartaric Acid has been suggested as 10 in 1,000,000, see Tartaric Acid.

Hydrogen Sulphide—The aqueous solution (1-20) of the salt should not be affected by T.S. of Hydrogen Sulphide, *P.G.*, slightly acidulated with Hydrochloric Acid should not respond to the tinge limit test for heavy metals, *U.S.P.*

Potassium or Sodium Hydroxide—When heated with Sodium Hydroxide T.S. it should not evolve Ammonia, *P.G.* The *U.S.I.* uses Potassium Hydroxide T.S.

Barium Nitrate—An aqueous solution (1-20) after the addition of Nitric Acid and the removal of the crystalline precipitate should not be affected by T.S. of Barium Nitrate, *P.G.*

Silver Nitrate—An aqueous solution (1-20) of the salt after treatment as in previous test should not be rendered more than opalescent by T.S. of Silver Nitrate, *P.G.*

Ammonium Oxalate—If 1 gramme of the salt be dissolved in 10 c.c. of Water and shaken with 5 c.c. of diluted Acetic Acid, the liquid poured off from the crystalline precipitate which separates out and diluted with an equal part of Water should not be affected within 1 minute by 8 drops of T.S. of Ammonium Oxalate, *P.G.*

Volumetric Determination—If 1 gramme of the salt be thoroughly ignited at red heat, and the residue extracted with boiling Distilled Water until the washings cease to react with Methyl Orange T S, the mixed filtrate and washings should require for complete neutralisation not less than 14 c.c. of Semi-normal Volumetric Solution of Hydrochloric Acid, Methyl Orange T S being used as indicator, *U S P*

The residue from the ignition of 1 gramme of the salt dissolved in Water, should require for exact neutralisation not less than 7 c.c. of Volumetric Solution of Sulphuric Acid, *B P*

Preparation

PULVIS SODÆ TARTARATÆ EFFERVESCENS. EFFERVESCENT TARTARATED SODA POWDER Commonly known as Seidlitz Powder *NO Syn*—PULVIS AEROPHORUS LAXANS, PULVIS EFFERVESCENS LAXANS

Sodium Potassium Tartrate, in dry powder, 120 grains, Sodium Bicarbonate, in dry powder, 40 grains Mix Wrap in blue paper Tartaric Acid, in dry powder, 38 grains Wrap in white paper

Dose—The quantities given above are intended for one dose The powder in blue paper is first dissolved in about half a pint of Water, and the powder in white paper added to it and the whole taken during effervescence

The chief Continental Pharmacopœias have a simple Effervescent Powder, made with Sodium Bicarbonate and Tartaric Acid, and also a compound powder containing similar ingredients to the above

Official in all the Foreign Pharmacopœias except Fr and Ital
Fr has Eau Saline Purgative gazeuse (Eau dite de Seidlitz), *see p 755*

Not Official

SODII ACETAS

SODIUM ACETATE

$\text{NaC}_2\text{H}_3\text{O}_2 \cdot 3\text{H}_2\text{O}$, eq 135.10

Fr, ACÉTATE DE SODIUM, Ger, Natriumacetat, Ital, ACETATO DI SODIO, Span, ACETATO SÓDICO

Colourless, translucent, monoclinic prisms, or as a white, granular, crystalline powder, possessing a saline, bitter taste The crystals are efflorescent in warm air, and should be kept in well-closed bottles in a cool place

Sodium Acetate is not official in the *B P*, but is official in both the *U S P* and the *P G*

Solubility—1 in 1 of Water, 1 in 30 of Alcohol (90 p.c.)

It has been employed as a diuretic in place of the Potassium salt, but is rarely used medicinally Used in the preparation of Acetic Ether

Foreign Pharmacopœias—Official in Fr, Ger, Hung, Ital, Jap, Mex, Russ, Swed, Swiss and U S

Tests—Sodium Acetate liquefies when heated and loses its Water of crystallisation, equivalent to 39.7 p.c. The *U S P* gives the liquefying point at 60° C (140° F), and states that at 123° C (253.4° F) it becomes dry and anhydrous When still more strongly heated it is decomposed, evolving empyreumatic vapours and leaving a black residue, which when dissolved in Water possesses a strong alkaline reaction towards red Litmus paper, and which effervesces on the addition of diluted acid The salt dissolves readily in Water, forming a colourless solution which is alkaline in reaction towards red Litmus paper but which produces little or no coloration with Phenolphthalein Solution The solution answers the tests

distinctive of Sodium given under that heading, and the addition of Ferric Chloride T S produces a deep red coloration, the solution on boiling yielding a brown flocculent precipitate of basic Ferric Acetate. When warmed with Sulphuric Acid it evolves a strong acetous odour, and when warmed with Sulphuric Acid and a small quantity of Alcohol (90 p c) it yields the characteristic odour of Ethyl Acetate (Acetic Ether), a minute quantity of the anhydrous salt when heated with a correspondingly minute quantity of Anomalous Anhydride yields a characteristic and highly poisonous odour of Cacodyl Oxide. The *U S P* requires it to contain at least 99.5 p c of pure Sodium Acetate as volumetrically determined by the process described below. The *P G* does not state what percentage of pure Sodium Acetate the salt should contain, nor does it give a method of determination.

The more generally occurring impurities, viz Arsenic, Lead, Copper, Iron and Zinc, Calcium, Potassium, Chlorides and Sulphates. Arsenic, if present, may be detected by the modified Gutzkow's test, Lead, Copper, Iron and Zinc, if present, may be detected by Hydrogen Sulphide, either in a solution rendered faintly acid by diluted Hydrochloric Acid or in a solution rendered alkaline with Ammonia. Calcium, if present, by the addition of Ammonium Oxalate Solution to an aqueous solution of the salt. Potassium, if present, by the turbidity produced on the addition of Sodium Bitartrate T S to a saturated aqueous solution of the salt. A 10 p c aqueous solution should not be rendered turbid by the addition of Barium Nitrate Solution, nor when acidified with Nitric Acid by Silver Nitrate Solution. The *P G* includes a separate test for the presence of Iron, requiring that 20 c c of a 1-20 aqueous solution should not assume a blue coloration on the addition of 0.5 c c of Potassium Ferriocyanide solution (1-20).

Volumetric Determination—A weighed quantity of 1 gramm of the salt is thoroughly carbonised at a temperature not exceeding a red heat, the residue is treated with boiling Water, the solution filtered, and the extraction of the residue continued with boiling Distilled Water until the washings no longer produce an alkaline reaction with Methyl Orange T S. The mixed filtrate and washings should require for complete neutralisation not less than 14.7 (14.74) c c of Semi-normal Volumetric Sulphuric Acid Solution, Methyl Orange T S being employed as an indicator of neutrality, *U S P*.

SODII THEOBROMINÆ ACETAS (Agurin) $C_7H_5N_3O Na C H_3O Na$, eq 282.13.—A white, crystalline powder, possessing a somewhat bitter, saline taste. It is soluble in Water. Introduced as a diuretic.

It should be kept in well closed glass bottles and protected as far as possible from exposure to the air, as it has a tendency to absorb Carbonic Anhydride.

It contains theoretically 63.8 p c of pure Theobromine, and shows a Theobromine content 10 p c in excess of that contained in Theobromine Sodium Salicylate.

Dose—3 to 8 grains = 0.2 to 0.52 gramme, given in form of a cachet, or a suspension with mucilage.

Tests—Theobromine Sodium Acetate dissolves in Water, forming a solution which is slightly alkaline in reaction towards red Litmus paper. The aqueous solution when neutralised and diluted with Hydrochloric Acid throws down a precipitate of Theobromine, which dissolves when shaken with Chloroform. If the chloroformic solution be separated and a few drops be evaporated to dryness on a water bath on a watch glass, the residue when treated with Chlorine Water and again evaporated to dryness on the water bath leaves a reddish brown residue which, when moistened with Ammonia Solution, affords a purple violet coloration. An aqueous solution of the salt affords with Ferric Chloride T S a deep red coloration, which when boiled affords a brownish red precipitate of basic Ferric Acetate. When heated with a small quantity of Sulphuric Acid and a little Alcohol (90 p c) it evolves a characteristic odour of Ethyl Acetate (Acetic Ether), the salt should leave no weighable residue when ignited with free access of air.

SODII ARSENAS.

ARSENIALE OF SODIUM (HYDROUS), *B P* '85 Na_2HAsO_4 , eq 184.78

FR, ARSENIATE OF SODIUM OFFICINAL, GER, NATHUMARSENIAL, ITAL, ARSENIATO BISODICO, SPAN, ARSENIATO SODICO

The anhydrous salt, Di-sodium Hydrogen Arsenate

An odourless, white, granular, amorphous powder

It should be kept in well-stoppered glass bottles and protected as far as possible from exposure to a moist atmosphere

The anhydrous salt, Di-sodium Hydrogen Arsenate, is official in the *B P*, and is officially directed to be prepared by dehydrating the crystallised Di-sodium Hydrogen Arsenate at a temperature of 148.9°C (300°F). Crystallised Di-sodium Hydrogen Arsenate is prepared by crystallising from Water the product resulting from the fusion of Arsenious Anhydride with a mixture of Sodium Nitrate and Sodium Carbonate. Sodium Arsenate (*U S P*) is the Di-sodium-ortho-arsenate containing 7 molecules of Water of crystallisation. Exsiccated Sodium Arsenate (*U S P*) is the anhydrous or almost anhydrous Di-sodium-ortho-arsenate. The salt is not official in the *P G*. The *Brussels Conference* has adopted the crystallised salt containing 7 molecules of Water of crystallisation.

The crystallised salt occurs as colourless, odourless, translucent, prismatic crystals, with a slightly alkaline reaction, having the formula $\text{Na}_2\text{HAsO}_4 \cdot 7\text{H}_2\text{O}$, eq 309.94. The *B P* title 'Arsenate of Sodium' (hydrous), *B P* '85, might very well have been omitted, as it is very misleading, and apt to give rise to serious error if the terms are taken to be synonymous. Some mention of the equivalents of the two salts might at least have been made under this heading: 1 gram of the anhydrous salt = 1.67 grains of the crystallised ($7\text{H}_2\text{O}$) salt, not 1.77 grains as given in the *B P* monograph on *Liquor Sodii Arsenatis*.

Solubility—1 in 4 of Water**Medicinal Properties**—Similar to those of *Liquor Arsenicalis*. See *ACIDUM ARSENIOSUM*, p 14**Dose.**— $\frac{1}{40}$ to $\frac{1}{10}$ grain = 0.0016 to 0.0065 gramme*Ital* maximum single dose of the crystals ($7\text{H}_2\text{O}$), 0.004 gramme maximum daily dose, 0.015 gramme**Prescribing Notes**—Generally employed in the form of the *Liquor*, may also be given in pills well triturated with Milk Sugar and 'Diluted Glucose,' q s**Official Preparation**—*Liquor Sodii Arsenatis***Antidotes**—See *ACIDUM ARSENIOSUM*, p 15**Foreign Pharmacopœias**—Official in Belg, dried salt, Dutch, Fl, Ital, Mex, Port, Span and Swiss, crystallised, Ital and U S both**Tests.**—Sodium Arsenate dissolves readily in Water, forming a solution which possesses an alkaline reaction toward red Litmus paper. It answers the tests distinctive of Sodium given under that heading. The 5 p.c. aqueous solution affords with Barium or Calcium Chloride Solutions a white precipitate soluble in Nitric Acid, with Silver Ammonio-Nitrate Solution a dark red precipitate soluble in Nitric Acid, with Magnesium Ammonio-Sulphate Solution a white

crystalline precipitate soluble in a diluted mineral acid. It is officially required to contain 99.8 per cent of pure anhydrous Sodium Arsenate, as gravimetrically determined by precipitating a solution of a weighed quantity of 1 gramme of the salt and 1 (presumably gramme) of Glacial Acetic Acid in 50 c.c. of Water with Lead Acetate, the *B.P.* mentions that such a solution should require 2.03 grammes of the Lead salt for complete precipitation. The Report of the Committee of Reference in Pharmacy states that the Lead Acetate test, which has been several times the subject of discussion, is substantially correct if carried out as described in an Acid solution. The official volume does not state whether the 1 of Glacial Acetic Acid is by weight or by measure. The *U.S.P.*, although requiring that the salt should contain not less than 98 per cent of anhydrous Disodium orthoarsenate, does not describe a method of determination. A volumetric process would have been preferable. A convenient method has been suggested (*Y.B.P.* '02, 505), the process recommends titrating a weighed quantity of the salt with Normal Volumetric Sulphuric Acid, using Methyl Orange Solution as an indicator of neutrality. Owing to the high molecular equivalent of the salt a good quantity should be employed for the determination, not less than 3 grammes of the crystallised or its equivalent of the anhydrous salt has been suggested.

The more generally occurring impurities are Lead, Copper and Iron, Aluminium, Calcium, Carbonates, Chlorides, Nitrates and Sulphates, a certain amount of moisture may also be present. 5 c.c. of a 1 in 20 aqueous solution of the salt when mixed with 1 c.c. of Ammonium Sulphide T.S. should not afford a dark coloration, indicating the absence of Lead, Copper and Iron. The aqueous solution when boiled with Ammonia Solution should not afford a white flocculent precipitate, indicating the absence of Aluminium, the aqueous solution should not yield on the addition of Ammonium Oxalate a white cloudiness or turbidity, indicating the absence of Calcium. The salt should not effervesce on the addition of a diluted mineral acid, indicating the absence of Carbonates. The aqueous solution when rendered acid with Nitric Acid should yield no pronounced turbidity on the addition of Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. When in aqueous solution of the salt is treated with Sulphuric Acid, the mixture being kept cool, it should yield no brown ring at the point of junction of the two liquids when Ferric Sulphate Solution is poured upon the surface, indicating the absence of Nitrates. Arsenite is sometimes present as an impurity, and may be detected by the Silver Nitrate test given below. The *B.P.* requires that it should not lose weight on being heated to 148.9° C (300° F), indicating the absence of Hydrous Sodium Arsenate. The *U.S.P.* requires that it should not lose weight when heated to 150° C (302° F), the *B.P.* includes tests for Magnesium, Potassium and Ammonium.

Silver Nitrate—If to 2 c.c. of an aqueous solution (1 in 20), 5 c.c. of Tenth normal Silver Nitrate Volumetric Solution be added, and the precipitate redissolved by a slight excess of Ammonia Water, no black precipitate of reduced Silver should appear on boiling (absence of Arsenite), *U.S.P.*,

Preparation

LIQUOR SODII ARSENATIS. SOLUTION OF SODIUM ARSEFATE
Sodium Arsenate, recently rendered anhydrous, $17\frac{1}{2}$ grains,
Distilled Water, *q s* to yield 4 fl oz (1 in 100)

After being made, this solution deposits a little Silica introduced in the preparation of the Arsenate, but, if filtered after a few days, remains clear.

It is about half the strength of Liquor Arsenicalis in Arsenic, as that preparation contains 1 p c of Arsenious Acid, and this 1 p c of Sodium Arsenate another difference is that Liquor Arsenicalis contains an Arsenite, and this an Arsenate

Dose.—2 to 8 minims = 0.12 to 0.5 gramme

11 minims contain $\frac{1}{10}$ grain of the anhydrous salt

Ph Ger maximum single dose, 0.5 gramme, maximum daily dose, 1.5 grammes, of the Potassium Arsenite Solution

Foreign Pharmacopœias—Official in U S, same as Brit, Dan, Port and Swiss, 1 in 500, Ital and Mex. have Solucion Arsenical de Pearson, 1 in 600

Tests.—Solution of Sodium Arsenate has a sp gr of 1.010 to 1.015. It is officially required to contain 1 p c w/v of anhydrous Sodium Arsenate, but no method is given by which this requisite preparation may be ensured. The *USP* requires that it shall contain 1 p c Arsenate corresponding in amount to not less than 1 p c of exsiccated Sodium Arsenate, but like the *BP* gives no method by which this proportion may be ensured. It should respond to the tests distinctive of Sodium Arsenate given under Sodii Arsenas, and should be free from the impurities there mentioned

SODII BENZOAS.

SODIUM BENZOATE

C_6H_5COONa , eq 143.01.

FR, BENZOATE DE SODIUM, GER, NATRIUMBENZOAT, ITAL, BENZOATO DI SODIO, SPAN, BENZOATO SODICO

A white, odourless, amorphous powder or having a faint odour of Benzoin when made from resin-sublimed Acid. It is obtained by neutralising Benzoic Acid with Sodium Carbonate.

The salt is official in the *USP*, but not in the *PG*. The *USP* requires that it shall contain not less than 99 p c of pure Sodium Benzoate.

It should be kept in well-stoppered glass bottles and in a cool atmosphere.

Solubility—1 in 2 of Water, 1 in 25 of Alcohol (90 p c)

Medicinal Properties.—Similar to Benzoic Acid, but less irritating, given in chronic cystitis in which there is alkaline and decomposing urine.

The Royal College of Physicians of London recommended a solution of 120 grains in a quart of hot Water injected into the bowel in cholera, if much pain, 15 to 30 minims of Laudanum may be added.—*L*, 92, 11 633.

Dose —5 to 30 grains = 0.32 to 2 grammes

Prescribing Notes —*May be given in cachets but generally employed as solution*

Incompatibles —Ferric Salts, Citric and Tartaric Acids, and Mineral Acids.

Foreign Pharmacopœias —Official in all except Dan., Ger., Norw. and Swed. Dutch has also Benzoyl Natrium cum Coffeino, Sodium Benzoate and Caffeine equal parts

Tests —Sodium Benzoate when heated melts, evolving an odour of Benzoic Acid, finally charring when strongly heated and leaving a residue which, when dissolved in Water, yields a solution having a strongly alkaline reaction towards red Litmus paper. The salt dissolves readily in Water, forming a colourless solution which possesses a faintly alkaline reaction towards red Litmus paper. The *USP* states that the aqueous solution is neutral or slightly alkaline towards Litmus paper. After separation of the Benzoic Acid it answers the tests distinctive of Sodium given under that heading. The aqueous solution of the salt affords with Ferric Chloride T.S. a buff coloured precipitate. A concentrated aqueous solution of the salt yields on the addition of sufficient Diluted Sulphuric Acid a bulky white crystalline precipitate, which when separated, washed till free from mineral acid and carefully dried should possess the m.p. and answer the tests distinctive of Benzoic Acid described under Acidum Benzoicum. It is officially required to yield from 97.2 to 98.7 p.c. of pure Sodium Benzoate as volumetrically determined by the process described below under the heading of Volumetric Determination. Three commercial samples contained an average of 4 p.c. of Water, which the volumetric test requiring 97 p.c. of anhydrous Sodium Benzoate does not recognise. The *USP* requires that it should contain not less than 99 p.c. of pure Sodium Benzoate as volumetrically determined by the process also mentioned in small type. It may be noticed that the Pharmacopœia leaves the choice of indicator to the operator. The *USP* states that Methyl Orange T.S. should be used as an indicator of neutrality. In carrying out the Pharmacopœia process of igniting a gramme of the salt, it will be found that considerable difficulty is experienced in burning off the carbonaceous matter, and the result is liable to be below the truth, owing to loss or incomplete washing of the partially incinerated residue. The method of direct titration similar to that given under Lithii Benzoas may be employed. 1 c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution is equivalent to 0.014301 gramme of Sodium Benzoate.

The more generally occurring impurities are heavy metals, Lead, Copper and Iron, Calcium, Ammonium, Carbonates, Chlorides or Sulphates. If the aqueous solution of the salt be slightly acidified with Diluted Hydrochloric Acid, and if the precipitated Benzoic Acid be separated by filtration, the filtrate should yield no darkening in colour when tested with Hydrogen Sulphide, either in the acidified filtrate as it stands, or after being rendered alkaline by the addition of Ammonia Solution, indicating the absence of Lead, Copper and Iron. The aqueous solution should afford no opalescence on the addition of Ammonium Oxalate Solution, indicating the absence

of Calcium. When boiled with Potassium or Sodium Hydroxide Solution it shall not yield an ammoniacal odour, nor shall the issuing vapour possess an alkaline reaction towards moistened red Litmus paper, indicating the absence of Ammonium salts. The aqueous solution should not yield an effervescence on the addition of Diluted Sulphuric Acid, indicating the absence of Carbonate. If the aqueous solution be acidified with diluted Nitric Acid and filtered from the precipitate of Benzoic Acid, the filtrate should yield only the faintest turbidity with Silver Nitrate or Barium Chloride Solutions, indicating the absence of more than traces of Chlorides and Sulphates. The *B P* also includes tests for Magnesium and Potassium.

Volumetric Determination—If 1 gramme of the dry salt be thoroughly ignited at red heat, and the residue extracted with boiling Distilled Water, until the washings cease to react with Methyl Orange T S, the mixed filtrate and washings should require for complete neutralisation not less than 13.85 c.c. of Semi-normal Hydrochloric Acid Volumetric Solution, Methyl Orange T S being used as indicator, *U S P*.

The residue from the ignition of 1 gramme of the salt, when dissolved in Water should require for neutralisation 6.8 to 6.9 c.c. of the Volumetric Solution of Sulphuric Acid, *B P*.

SODII BICARBONAS.

SODIUM BICARBONATE

NaHCO₃, eq. 83.43

FR., CARBONATE ACIDE DE SODIUM, GER., NATRIUMBICARBONAT,
ITAL., BICARBONATO DI SODIO, SPAN., BICARBONATO SODICO

Small, opaque, prismatic crystals, or an odourless, white, micro-crystalline powder, possessing a saline taste and alkaline reaction.

It should be kept in well-closed vessels and in a cool atmosphere.

Solubility—1 in 12 of Water, insoluble in Alcohol (90 p.c.)

Medicinal Properties—Analogous to those of Potassium Bicarbonate, but it is much more frequently given, as it is only feebly depressant and is more slowly absorbed than the Potassium salt. Employed as a gastric sedative half an hour before food, and as an antacid in preventing the eructations and pain of hyperacidity half an hour after food. In the Uric Acid diathesis the salts of Potassium and Lithium are preferable, as they form more soluble salts with Uric Acid. Very large doses are given in the acid intoxication and coma of diabetes. Moistened with Water, it is an excellent remedy to the stings of wasps and gnats, a lotion relieves itching.

Alkalis are beneficial in dyspepsia when given half an hour before food, not, as is sometimes taught, by increasing gastric secretion, but by inhibiting it, and so for the time being resting the stomach—(W. E. Dixon) *B M J* '06, ii 1459. Large doses the most successful treatment of acetonuria—*L* '07, i 511.

Sodium salts accelerate the conversion of gelatinous Sodium Bi-urate into the crystalline variety, and their employment in the treatment of gout is apparently not desirable—*L* '00, i 931, *B M J* '00, i 836.

5 cachets daily, each cachet containing 30 grains, increasing the dose if necessary, in the vomiting of pregnancy—*Pr* lxvii 244.

Large doses (100 to 125 grains in 24 hours) in recurring vomiting of infancy
—*M A '04*, 379

A sterile solution injected through a Eustachian catheter is probably the most efficacious treatment of non suppurative middle ear disease —*B M J '04*, 11206

120 to 180 grains per day in the recurrent vomiting of childhood —*B M J '04*, 11350, 351

Dose — 5 to 30 grains = 0.32 to 2 grammes

Prescribing Notes — *May be prescribed in cachets, powders, or in solution. It is also given in Compressed Tablets*

20 of Sodium Bicarbonate are neutralised by 16.7 of Citric Acid, and by 17.8 of Tartaric Acid

Official Preparation — *Trochiscus Sodii Bicarbonatis*. Used in the preparation of *Caffeine Citras Effervescent*, *Ferrum Arsenas*, *Ferrum Phosphoricum*, *Lithium Citras Effervescent*, *Magnesium Sulphas Effervescent Pulvis Sodici Tartaratus Effervescent*, *Sodii Citro Tartarus Effervescent*, *Sodii Phosphas Effervescent*, *Sodii Sulphas Effervescent*, *Spiritus Aethenis Compositus*, and 'Soluble Saccharum'

Not Official — *Sal Emsanum Factitium*, *Trochisci Bicarbonatis Natrii Compositi*, *Collunarum Alkalinum*, *Collunarum Alkalinum Co.*, *Mistura Sodici Composita*, *Nebula Alkalina*, and *Nebula Alkalina Composita*

Foreign Pharmacopœias — Official in all

Tests — Sodium Bicarbonate yields the tests distinctive of Sodium given under that heading. When heated it loses Carbon Dioxide and Water, being converted into Sodium Carbonate. It loses at a temperature of 100° C (212° F) (according to the *U S P*) about 36.5 p.c of its weight. According to the *I'G*, 100 parts of the salt previously dried over Sulphuric Acid shall leave, after ignition at a dull red heat, not more than 63.8 parts by weight of residue, corresponding to a loss of not less than 36.2 p.c. It effervesces strongly on the addition of a diluted mineral acid, yielding a colourless and odourless gas which, when passed into Lime Water, yields a white precipitate soluble in a sufficient excess of the gas or in diluted mineral acids. The salt is soluble in Water, forming a colourless solution which is alkaline in reaction towards red Litmus paper, it yields no precipitate with Magnesium Sulphate Solution. It is officially required to indicate 98.4 to 99.3 p.c of pure Sodium Bicarbonate as volumetrically determined by titrating 1 gramme of the salt with Volumetric Sulphuric Acid Solution, from 11.8 to 11.9 c.c should be necessary, the choice of an indicator of neutrality is left to the operator. The *U S P* requires that it should contain not less than 99 p.c of pure Sodium Bicarbonate as volumetrically determined by titrating 2 grammes of the salt with Normal Volumetric Sulphuric Acid Solution, by the method indicated in small type below under the heading of Volumetric Determination. The *U S P* directs the use of Methyl Orange Solution as an indicator of neutrality.

The more generally occurring impurities are Lead, Copper, Iron, Aluminium, Calcium, Ammonium, Chlorides, Sulphates, Thiocyanates, and Sodium Carbonate. The solution of the salt should afford no darkening in colour, either in acid or in alkaline solution, on the addition of Hydrogen Sulphide, indicating the absence of Copper,

Lead, and Iron Standards have been suggested (*CD* '08, 1. 796) of 5 parts per million for Lead, and 2 parts per million for Arsenic. A standard solution for Chloride is 0.1 per cent, calculated as Sodium Chloride. When dissolved in diluted Hydrochloric Acid, Ammonium Chloride added, and Ammonia Solution in faint excess, it shall yield no flocculent precipitate, indicating the absence of Aluminium. It should yield no opalescence on the addition of Ammonium Oxalate Solution to an aqueous solution slightly acidified with Acetic Acid, indicating the absence of Calcium. When heated in a dry test-tube it should not evolve an ammoniacal odour, nor should the vapour emitted possess an alkaline reaction towards moistened red Litmus paper. When the aqueous solution is acidified with Nitric Acid it should answer the test given under the heading of Silver Nitrate given below, and when the aqueous solution is saturated with Acetic Acid it should respond to the Barium Nitrate test described below, indicating the absence of Chlorides and Sulphates. The absence of Thiocyanates may be determined by the under-mentioned test with Ferric Chloride. The *BP* employs Mercuric Chloride T.S. as a means of distinguishing Sodium Carbonate from Sodium Bicarbonate. The test has been discarded by both the *USP* and the *PG*. The *BP* requires that a solution of the salt in cold Water should give, with Mercuric Chloride T.S., a whitish precipitate, becoming brownish-red on standing, soluble Carbonates being stated in the Appendix to afford a brownish-red precipitate with Mercuric Chloride T.S. Howard has pointed out (*CD* '98, 1. 675) that a pure sample will not pass the *BP* tests. Attfield (*Digest of Researches and Criticisms Report for 1898*) replies that the test is clearly not given as a 'pass' test of purity, but only as a 'distinction' test, and suggests that the critic missed an opportunity of recommending the addition of the following useful words to the official sentence: 'A solution of the salt in cold Water gives either no precipitate immediately or only a whitish precipitate, becoming reddish-brown on standing'. The *USP* and *PG* adopt practically the same method for determining the presence of the Normal Carbonate, which is described under the heading of Phenolphthalein in small type below. Traces of Sodium Carbonate, and also of Water, are probably present in all commercial Sodium Bicarbonates, but it may still pass the *BP* titration test, owing to the counterbalancing influence of the two impurities. The actual Carbonate may be estimated by adding an excess of Normal Volumetric Sodium Hydroxide Solution free from Carbonate, then an excess of Barium Chloride Solution, and titrating with Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. It has been recommended that the tests for Carbonate in the present official volume should be replaced by one on the lines of that in the *USP*, in the next revision. The *BP* includes, in addition to the above list of substances, Magnesium, Potassium, Sulphites and Thiosulphates as likely impurities, and requires that the salt should yield the customary no characteristic reaction with the tests for the three former and gives a test with Ferric Chloride Solution for the

detection of the last named, no red coloration should be produced when the reagent is added to an aqueous solution acidified with Hydrochloric Acid

Phenolphthalein—A solution of 1 gramme of the salt in 20 c.c. of Water dissolved without agitation at a temperature not exceeding, 15° C (59° F) should not be immediately coloured red by 3 drops of Phenolphthalein T S, and any faint reddening produced should be discharged by 0.2 c.c. of Normal Volumetric Hydrochloric Acid Solution, *P G*. The *U S P* states that in such a solution a red tint should not be produced immediately on the addition of 0.2 c.c. of Normal Volumetric Hydrochloric Acid Solution and 2 drops of Phenolphthalein T S

Ferric Chloride—5 c.c. of an aqueous solution (1 in 20) should not be coloured red by 1 drop of T S of Ferric Chloride, *U S P*, also in *L P* and *P G*, the *B P* does not give quantities, and the *P G* uses a 1-50 solution acidulated with Nitric Acid

Hydrogen Sulphide—An aqueous solution of the salt (1-50) saturated with Acetic Acid should not be affected by T S of Hydrogen Sulphide, *P G*. An aqueous solution (1-20) acidulated with Hydrochloric Acid should not respond to time limit test for heavy metals, *U S P*

Barium Nitrate—An aqueous solution of the salt (1-50) saturated with Acetic Acid should be rendered not more than faintly opalescent by Barium Nitrate T S within 2 minutes, *P G*

Silver Nitrate—An aqueous solution (1-50) acidulated with Nitric Acid should be clear and should not show more than a whitish opalescence within 10 minutes on the addition of T S of Silver Nitrate, *P G*

Volumetric Determination—23.7 (23.74) c.c. of Normal Sulphuric Acid Volumetric should be necessary to completely neutralise 2 grammes of the salt, Methyl Orange T S being used as indicator, *U S P*

Preparations

TROCHISCUS SODII BICARBONATIS—SODIUM BICARBONATE
LOZENGE

Contain 3 grains in each, with Rose basis

Dose—1 to 6 lozenges

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Ital, Jap, Mex, Norw, Port, Russ, Spm, Swiss and U S

SODII CITRO-TARTRAS EFFERVESCENS EFFERVESCENT
SODIUM CITRO-TARTRATE

Sodium Bicarbonate, 51, Tartaric Acid, 27, Citric Acid, 18
Refined Sugar, 15, all in powder, made into granules, the yield of which is about 100

Dose—60 to 120 grains = 4 to 8 grammes, as a mild, saline purgative

Not Official

SAL EMSANUM FACTICIUM—Dried Sodium Sulphite, 7, Potassium Sulphate, 13, Sodium Chloride, 325, Sodium Bicarbonate, 655—*Dutch*

TROCHISCI BICARBONATIS NATRICI COMPOSITI—Salis Emسانی faction, 25, Sugar, 75—*Dutch*

COLLUNARIUM ALKALINUM—Sodium Bicarbonate and Borax, of each 3 grains, Phenol, 1 grain, White Sugar, 5 grains, Water, to 1 oz—*Throat*

COLLUNARIUM ALKALINUM CO—Sodium Bicarbonate, Borax, Sodium Chloride, of each 2 grains, White Sugar, 5 grains, Water, to 1 oz—*Throat*

MISTURA SODÆ COMPOSITA—Gentian Root, 5 grains, Rhubarb Root, 2 grains, Ginger, 1 grain, Sodium Bicarbonate, 10 grains, Peppermint Water, to 1 fl oz. Macerate the Gentian, Rhubarb and Ginger sliced, with the Sodium Bicarbonate in the Peppermint Water for 24 hours, then press out the liquor, strain, and pour Peppermint Water over the strainer until the product measures 1 fl oz.—*St Thomas's*

This has been incorporated in the *B P C* with the *syn* Peacock's Stomachic Mixture

NEBULA ALKALINA—Sodium Bicarbonate, 15 grains, Borax, 15 grains, Carbolic Acid, 4 grains, Glycerin, 45 minims, Water, to 1 oz.—*Throat*

Nebula Alkalina Composita—Sodium Bicarbonate, 1 50, Borax, 1 50, Carbolic Acid, 0 75, Glycerin, 25, Distilled Water, *q s* to produce 100—*B P C*

SODII BROMIDUM.

SODIUM BROMIDE

NaBr, eq 102 23

FR, BROMURE DE SODIUM, GER, NATRIUMBROMID, ITAL, BROMURO DI SODIO, SPAN, BROMURO SODICO

Minute white crystals, or a white crystalline powder, possessing a saline, slightly bitter taste. It may be prepared in a similar manner to Potassium Bromide, employing Sodium Hydroxide in place of Potassium Hydroxide.

As this salt is very deliquescent it should be kept in well-stoppered bottles. It may be prepared either anhydrous, or containing 2H₂O.

Solubility.—5 in 6 of Water, and measures 7½, 1 in 16 of Alcohol (90 p c)

Medicinal Properties.—Similar to Potassium Bromide, but less depressant, and more easily tolerated by the stomach.

It has been recommended as a remedy for sea sickness in 60 grain doses 3 times a day for at least 2 days before embarkation on a long voyage, the doses being reduced to half when on board.—*B M J* '81, 11 730

Deprivation of salt and substitution of Bromide, about ½ oz per week being taken in epilepsy.—*B M J* '03, 1 552

A nightly draught containing from 20 to 30 grains, together with a cachet containing 10 grains of Chloralamide, and followed by a second cachet containing 10 grains of Chloralamide if sleeplessness persists, in the treatment of insomnia accompanying the rapid heart of influenza.—*J* '99, 11 1079

In the treatment of acute mania, 2 drim in a half tumbler of Water every 2 hours until 1 oz is given the first day, a similar amount given on the second day, and this may suffice to effect the result desired, which is not at its height until the fourth or fifth day, ceasing the administration for 24 hours, when drowsiness is so profound that the patient cannot be roused, or if when roused talk is incoherent.—*B M J* '00, 1 134

Given in the treatment of the Morphine, Chloral and Cocaine habits. 30 grains twice daily increasing the dose to 40, 50, 60, and even 120 grains if required.—*T. G* '90, 600, 30 to 60 grains every 3 or 4 hours for a day or two.—*B M J* '97, 11 77, 120 grains in solution every 2 hours for the first 2 days, and 60 grains during the third day.—*B M J* '99, 1 898

Dose.—5 to 30 grains = 0 32 to 2 grammes

Prescribing Notes.—Generally given in solution, it may be prescribed in powders if carefully wrapped in Tin foil. It is also given in Compressed Tablets and Effervescent Granules.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Norw, Russ, Span, Swed, Swiss and U S

Tests—Sodium Bromide when heated melts, and when strongly heated volatilises slowly without decomposition. It answers the tests distinctive of Sodium given under that heading. It dissolves readily in Water, forming a clear solution which is neutral to Litmus, or only faintly alkaline in reaction towards red Litmus paper. The aqueous solution yields, on the addition of Silver Nitrate Solution, a yellowish curdy precipitate, practically insoluble in Ammonia Solution, insoluble in Nitric Acid, readily soluble in Potassium Cyanide Solution. With Chlorine Solution it affords a reddish coloration, which passes into chloroformic solution when shaken with that menstruum. A small portion of the salt, heated with Sulphuric Acid and a little Manganese Dioxide, evolves reddish vapours of Bromine, which impart an orange yellow colour to Starch paper. It is officially required, when dry, to contain not less than 97.9 nor more than 99.98 pc of pure Sodium Bromide, as volumetrically determined by titration with Tenth normal Volumetric Silver Nitrate Solution, as indicated below under the heading Volumetric Determination. The *U S P* requires that it shall contain, when dried, not less than 97 pc of pure Sodium Bromide, as volumetrically determined by the method given in small type below. The *Pf* requires that the salt dried at 100° C (212° F) shall contain not more than 100.6 pc of pure Sodium Bromide, as volumetrically determined by the process also given below.

The more generally occurring impurities are Arsenic, Lead, Copper, Iron and Zinc, Barium and Calcium, Ammonium, Carbonates, Cyanides, Bromates, Iodates, Chlorides, Iodides, Sulphates and Thiocyanates. Arsenic, Lead, Copper, Iron and Zinc may be detected by Hydrogen Sulphide, either in a solution rendered slightly acid or in a solution rendered faintly alkaline by Ammonia Solution. Barium may be detected by the test described below under the heading of Potassium Sulphate. The aqueous solution should afford no opalescence when tested with Ammonium Oxalate Solution, indicating the absence of Calcium. When boiled with Potassium Hydroxide Solution it should afford no ammoniacal odour, nor should the evolved vapours exhibit an alkaline reaction towards moistened red Litmus paper, indicating the absence of Ammonium salts. An aqueous solution of the salt should yield no effervescence on the addition of Diluted Hydrochloric Acid, indicating the absence of Carbonates, nor should an odour of Hydrocyanic Acid be noticeable when the acidified solution is gently warmed, indicating the absence of Cyanides. Bromates, if present, may be detected by the Sulphuric Acid test given in small type. The aqueous solution, when mixed with Potassium Iodide Solution and Tartaric Acid, should not yield a blue coloration on the addition of Starch Mucilage, indicating the absence of Iodate. If the aqueous solution be completely precipitated with Silver Nitrate Solution and the precipitate be treated with Ammonia Solution and filtered, the filtrate shall yield only a faint turbidity when acidified with Diluted Nitric Acid,

indicating the absence of more than a trace of Chloride Iodides, if present, may be detected by the test described under the heading of Chlorine Water and Chloroform given in small type. The aqueous solution should not afford a distinct turbidity on the addition of Barium Chloride Solution. The *P G* includes a separate test for Iodine, which is described under the heading of Potassium Ferriocyanide in small type. The *BP* includes a test for Thiocyanates, requiring that Ferric Chloride T S should not cause a red coloration in an aqueous solution of the salt. It has been suggested (*P J* '01, 1 460) that the following modification of the Thiocyanate test should be made in the next revision of the *BP*. A weighed quantity of a gramme of the salt dissolved in 10 c c of Water should give a yellow and not a red or reddish-brown coloration on the addition of 2 drops of Ferric Chloride T S (absence of more than 0.01 p c of Ammonium Thiocyanate). Neither the *USP* nor the *P G* includes a test for Thiocyanate.

Sulphuric Acid—If diluted Sulphuric Acid be dropped upon some of the powdered salt no yellow colour should appear at once, *P G* and *USP*.

Hydrogen Sulphide—The aqueous solution of the salt (1-20) should not be affected by T S of Hydrogen Sulphide, *P G*. The *USP* requires that such an aqueous solution slightly acidulated with Hydrochloric Acid should not respond to the time-limit test for heavy metals.

Phenolphthalein—If 1 c c of 1 gramme of the salt in 10 c c of Water be acidulated with Sulphuric Acid Volumetric Solution added should not yield any colour with a drop of Phenolphthalein T S, even after boiling, *USP*.

Potassium Sulphate—10 c c of an aqueous solution (1-20) acidulated with Hydrochloric Acid should not be rendered turbid by 1 c c of T S of Potassium Sulphate, *USP*. The *P G* requires that an aqueous solution (1-20) should not be affected by diluted Sulphuric Acid.

Potassium Ferriocyanide—20 c c of an aqueous solution (1-20) previously acidulated with a few drops of Hydrochloric Acid should not be rendered blue by 0.5 c c of T S of Potassium Ferriocyanide, *P G*.

Chlorine Water and Chloroform—If to 10 c c of an aqueous solution of the salt (1 in 20) 1 c c of Chloroform be added, and then Chlorine Water which has been diluted with an equal volume of Water be cautiously introduced, drop by drop, with constant agitation, the liberated Bromine will dissolve in the Chloroform, imparting to it a yellow to orange colour, free from any violet tint, *USP*.

Volumetric Determination—Not less than 95.8 and not more than 97.8 c c of Silver Nitrate Volumetric Solution should be necessary for the complete precipitation of 1 gramme of the dry salt dissolved in Water, *BP*, 10 c c of an aqueous solution (3 grammes in 100 c c) of the salt, which has been dried at 100° C (212° F), should after the addition of a few drops of Potassium Chromate T S require not more than 29.3 c c of Tenth-normal Silver Nitrate Volumetric Solution to produce a red colour, *P G*, a solution of 0.3 gramme of dried salt in 50 c c of Water should require not more than 28.5 nor more than 30 c c of Tenth-normal Volumetric Silver Nitrate Solution to produce a permanent red colour, *USP*.

Rubidium Bromide in doses of 5 to 30 grains = 0.32 to 2 grammes, and **Rubidium Ammonium Bromide**, in doses of 10 to 40 grains have been introduced as substitutes for the alkaline Bromides in epilepsy.

Not Official

SODII CACODYLAS

 $\text{NaAs}(\text{CH}_3)_2\text{O}_2$, eq 158.96

White, odourless crystals, or as a white, amorphous, deliquescent powder.

It may be obtained by exactly neutralising Cacodylic Acid with the quantity of Sodium Hydroxide indicated by titration with Normal Volumetric Sodium Hydroxide Solution. It may be prepared anhydrous, but the anhydrous salt is very deliquescent, as it usually exists commercially it contains 2 to 3 molecules of Water of crystallisation.

It should be kept in well stoppered glass bottles and exposed as little as possible to the air, as it is of a deliquescent nature and readily absorbs moisture.

Solubility—2 in 1 of Water, 1 in 1 of Alcohol (90 p.c.)

It has been recommended on account of its lesser toxicity in all cases where Arsenic is usually employed, e.g., in tuberculous disease, in emu psoriasis and skin affections. It has also been used in the treatment of certain affections of the eye. When administered by the mouth in the form of pill or in solution it frequently imparts a disagreeable alliaceous odour to the breath, but when administered by hypodermic injection this objectionable feature is absent.

Professor Fraser has shown that when a salt of Cacodylic Acid is administered, it is absorbed and is eliminated, but the Arsenic it contains is so firmly combined that it does not become dissociated, and is therefore incapable of forming any compound in the body which can produce the well known pharmacological activities of the usual therapeutic compounds of Arsenic. It has been found by Crocker to be a failure in skin diseases.—*B M J* '02, i 712, '02, ii 656, *L* '02, i 748, '03, i 785.

General references.—*L* '00, ii 1446, 1923, '01, i 1462 '02, i 676, *B M J* '00, ii 1823, '01, i 120, *B M J* *L* '01 ii 32, 49, 83 *P J* '00, ii 721, '02, ii 336, 697, *C D* '02, i 59, 291, 466, *T G* '01, 790. Efficacious in yaws, *L* '07, ii 1459.

Foreign Pharmacopœias—Official in F.

Tests—Sodium Cacodylate melts at a moderately low temperature. It dissolves readily and completely in Water, forming a clear solution possessing a faintly alkaline reaction towards red Litmus paper. It yields no precipitate with Hydrogen Sulphide. It may be quantitatively determined by titration with Tenth normal Volumetric Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutrality. Cacodylic Acid is in itself neutral in reaction towards Methyl Orange Solution, whilst the Sodium Cacodylate is readily decomposed by the Volumetric Hydrochloric Acid Solution. 1 c.c. of Tenth normal Volumetric Hydrochloric Acid Solution corresponds to 0.015896 gramme of anhydrous Sodium Cacodylate. Occasionally commercial samples are found which contain free Cacodylic Acid, although there is no reason why, if carefully prepared, the resulting Cacodylate should not be neutral. In these cases, the free Cacodylic Acid may be determined by titration with Tenth normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 c.c. of Tenth normal Volumetric Sodium Hydroxide Solution is equivalent to 0.013708 gramme of pure Cacodylic Acid. The titration may then be continued with Tenth-normal Volumetric Hydrochloric Acid Solution, using Methyl Orange Solution as an indicator of neutrality. The number of c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution required to neutralise the free Cacodylic Acid should be deducted from the number of c.c. of Tenth normal Volumetric Hydrochloric Acid Solution required for the total titration with Methyl Orange Solution, the difference being calculated into anhydrous Sodium Cacodylate. 1 c.c. of Tenth normal Volumetric Hydrochloric Acid Solution corresponds to 0.015896 gramme of anhydrous Sodium Cacodylate. An aqueous solution of the salt when acidified with Nitric Acid should yield little or no turbidity on the addition of Silver Nitrate or of Barium Chloride Solution, indicating the absence of more than traces of Chlorides or Sulphates. It should not yield a precipitate on being rendered faintly alkaline with Lime Water, indicating the absence of Arsenic or Arsenious Acids and Oxalates. Cacodylates may be

distinguished from Methylarsenates and the presence of traces of the former in the latter detected by the reaction with an acid solution of Fe^{+3} . Dissolving 20 grammes of Sodium Hypophosphite in 20 c.c. of 200 c.c. of pure Hydrochloric Acid, a little Sodium Chloride crystallises out and may be separated by straining through absorbent Cotton-Wool. In applying the test 1 c.c. of a solution containing a trace of Cacodylate is added to 10 c.c. of the Acid Hypophosphite reagent, and the tube corked and allowed to remain at rest, an odour of Cacodyl will be developed after a time, even $\frac{1}{2}$ mg. of Sodium Cacodylate giving a perfectly distinct odour in 12 hours, but no precipitate of Arsenic. In solutions containing larger quantities of Cacodylate a deposit of Arsenic is slowly formed on the sides of the tube. In the case of Methylarsenates no odour is evolved, the whole of the Arsenic in combination is precipitated at once.

INJECTIO SODII CACODYLATIS—A sterilised solution, containing $\frac{1}{4}$ grain of pure Sodium Cacodylate in 17 minims. Also put up in glass capsules, each containing 1 c.c.

Elixir Sodii Cacodylatis—An elixir, each fl. oz. of which contains $\frac{1}{2}$ grain pure Sodium Cacodylate.

Globules Sodii Cacodylatis—Globules containing $\frac{1}{2}$ grain pure Sodium Cacodylate, also globules containing $\frac{1}{4}$ grain.

ACIDUM CACODYLICUM Cacodylic Acid $\text{HAS}(\text{CH}_3)_2\text{O}$, eq. 137.08

Tests—Cacodylic Acid or Di-methyl Arsenic Acid melts at about 200°C (392°F). It dissolves readily and completely in Water, forming a clear solution which possesses an acid reaction towards Litmus paper and towards Phenolphthalein Solution. The acid is reduced to Cacodyl Oxide by Phosphorous Acid, and is converted into Cacodyl Sulphide by Hydrogen Sulphide in the presence of Water, but by dry Hydrogen Sulphide it is converted into Thio-cacodylic Acid. The alcoholic solution gives a precipitate with Alcoholic Mercuric Chloride Solution. It may be determined quantitatively by titration with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 c.c. of Normal Volumetric Sodium Hydroxide Solution is equivalent to 0.13708 gramme of pure Cacodylic Acid. The aqueous solution when acidified with Nitric Acid should yield at the most but a faint turbidity with Silver Nitrate Solution, or with Barium Chloride Solution, indicating the absence of more than traces of Chlorides or Sulphates. When rendered faintly alkaline with Lime Water it should yield no decided turbidity or precipitate indicating the absence of Arsenious or Arsenic Acids and Oxalates. When used with free access of air it should burn leaving no weighable residue.

FERRI CACODYLAS (Iron Cacodylate)—A yellow, or reddish-yellow, amorphous powder, soluble 1 in 15 of Water, insoluble in Alcohol (90 p.c.). Successful in anæmia.—*B.M.J.* '02, i 713, *B.M.J.E.* '00, ii 58, '02, ii 57, *P.J.* '00, ii 724, '03, i 197.

Dose— $\frac{3}{4}$ to 5 grains by the mouth. 17 minims = 1 c.c. of the under-mentioned solution hypodermically.

Tests—Iron Cacodylate dissolves in Water, forming a clear solution which possesses an acid reaction towards blue Litmus paper. When ignited it leaves a residue consisting of Ferric Oxide which, when dissolved in Hydrochloric Acid containing a trace of Nitric Acid, yields the tests distinctive of Ferric salts given under Ferrum, p. 504. Commercial samples yield about 28.9 p.c. of Ferric Oxide on ignition. An alcoholic solution of Mercuric Chloride yields a yellow precipitate. The Iron Oxide should be also free from the impurities mentioned under Liquor Ferri Perchloridi Fortis, with the exception of Arsenic.

Injectio Ferri Cacodylatis—A sterilised solution containing $\frac{3}{4}$ grain of Iron Cacodylate in 17 minims of solution. Also a double strength solution containing $1\frac{1}{2}$ grains in 17 minims. Used with success in anæmia.

MAGNESII CACODYLAS (Magnesium Cacodylate)—A white, amorphous powder, readily soluble in Water. Used for the same purpose as the Sodium compound.—*P.J.* '02, i 123.

DI-SODII METHYLARSENAS Di sodium Methylarsenate $\text{Na}_2\text{CH}_3\text{AsO}_3$, $6\text{H}_2\text{O}$, eq 290.09—Colourless translucent crystals, or masses of crystals, or as a white granular powder. Soluble 1 in $1\frac{1}{2}$ of Water, insoluble in Alcohol (90 p.c.) Introduced as a comparatively non-toxic preparation of Arsenic, and employed in phthisis, in anæmias, and in malarial cachexia.—*L* '02, i 623 *B M J* '02, i 801, *B M J F* '02, i 68, *P T* '02, i 253, 256, 282

Dose— $\frac{1}{2}$ to 1 grain = 0.03 to 0.06 gramme in solution, hypodermically, or in pill form

Though containing much Arsenic, practically in inert substance, and even in enormous quantities it was incapable of producing the well-defined pharmacological action, and the well-recognised toxic effects of the Arsenic ion, and also incapable of exerting the remedial or therapeutic influences which were those of the older and commonly used compounds of Arsenic.—*B M J* '03, i 428, *L* '03, i 304

Non-toxic effect of organic Arsenic compounds demonstrated by Lawson sixty years ago.—*L* '03, i 174

Tests—Di sodium Methylarsenate dissolves readily in Water forming a clear solution possessing a strongly alkaline reaction towards red Litmus paper. It may be determined quantitatively by titration with Tenth normal Volumetric Sulphuric Acid, using Rosolic Acid Solution as an indicator of neutrality, in the event of free Methylarsenic Acid being present, it may be determined by first titrating with Tenth normal Volumetric Sodium Hydroxide Solution, using a similar solution as an indicator. 1 c.c. of Tenth normal Volumetric Sodium Hydroxide Solution represents 0.018905 gramme of Methylarsenic Acid, the titration may then be continued with Tenth normal Volumetric Acid Solution as above, deducting the number of c.c. of Tenth normal Volumetric Sodium Hydroxide Solution with Rosolic Acid Solution, the difference is calculated into terms of Di sodium Methylarsenate. 1 c.c. of Tenth normal Volumetric Sulphuric Acid Solution represents 0.029009 gramme of Di sodium Methylarsenate. It may also be determined by the much more intricate method of indirect Silver titration according to the method of Volhard. Di sodium Methylarsenate may be distinguished from Sodium Cacodylate and the presence of a trace of the latter in an excess of the former by the Acid Hypophosphite reagent mentioned under Sodium Cacodylate. In performing the test it is only necessary to dissolve 0.2 of a gramme of the salt in 10 c.c. of the reagent, cork the tube and allow the mixture to stand for 12 hours, in the presence of even $\frac{1}{2}$ mg. of Sodium Cacodylate a marked odour of Cacodyl will be evident.

The salt is also known under the commercial names of **Airhenal** and **Arsinyal**.

Injectio Arsinyal—A sterilised solution containing $\frac{1}{2}$ grain of the pure salt in 17 minims.

Elixir Arsinyal—An elixir containing $\frac{1}{2}$ grain of the pure salt in 1 fl. dram.

ATOXYL is a white, odourless, crystalline powder, soluble 1 in 5 of Water, insoluble in Alcohol (90 p.c.). When originally introduced, the composition of Atoxyl was stated to be Met-arsenic Acid Anilide, but on account of the indifferent solubility of the salt, the Sodium salt of Met-arsenic Acid Anilide was introduced. It is not now believed to be an Aniline compound at all, but the Sodium salt of Para-aminophenylarsonic Acid. The *B P C* mentions that Sodium Anilarsenate (*B P C*) is also known under the trade name Atoxyl, but this statement is incorrect.

Dose— $\frac{1}{4}$ to 3 grains = 0.05 to 0.2 gramme. **Globules**— $\frac{1}{4}$ grain (0.032 gramme) in each, **Elixir** = $\frac{1}{2}$ grain (0.032 gramme) in each fl. dram (3.6 c.c.), **Injection** = $\frac{1}{2}$ grain (0.032 gramme) in 17 minims (1.0 c.c.).

A reference to this interesting compound, and the results of its action on trypanosomes, is recorded.—*B M J* '05, i 1112. The general conclusion formulated is that treatment with this compound is in many ways superior to the ordinary arsenical treatment, on account of the quicker but prolonged action of the drug on the parasite, the large doses which can be given without toxic symptoms, and the entire absence of any tendency to sloughing.

It is best administered intravenously in high doses and for a long period, pushing it to the maximal amount that the patient can stand without headache and nausea, at the same time building up the patient in every way possible that will conduce to a lessening of the anæmia.

A further reference to its use in trypanosomiasis — *B M J* '06, i 1057 Inject subcutaneously in the form of a 20 p c solution in normal saline, in doses of 0.6 c c for 4 to 6 days, then increase the dose to 0.8 c c for 4 to 6 days, then to 1 c c per diem, continuing this dose until signs of intoxication begin to appear.

In trypanosomiasis, should be given as soon after infection as possible, and fresh solutions only must be used — *L* '08, i 113

Used with benefit in yaws (fiambœsia) both as a curative and preventive — *L* '07, ii 1459

Complete recovery from malaria after 2 injections of 1.2 c c of 10 p c solution — *B M J E* '07, ii 52

Leading article on its value in trypanosomiasis — *B M J* '07, ii 1738

In syphilis — *B M J* '07, i 1458, but with caution — *B M J* '07, ii 294

A specific in psoriasis — *L* '07, i 1151

Results in trypanosomiasis by the Atoxyl and Mercury method distinctly encouraging — *B M J* '07, ii 624, 685

It is stated to be at least 20 times less toxic than Arsenic, at any rate when given hypodermically. The solution of Atoxyl is readily decomposed by acids, alkalis, and light. It should therefore be made up fresh every few days. The solution turns yellow or brownish, and whereas a fresh solution when injected causes no irritation, the solution used after it has been kept some days causes inconvenience and irritation, the syringe used should be sterilised by boiling and not be placed in any antiseptic. The practice in London is to commence with 1 grain, repeating the injections every second day, adding $\frac{1}{2}$ grain more each time until a dose of 3 grains is reached, this being then continued. A few cases have had as much as 8 grains twice in a week. In Brussels the custom is to inject that quantity of fluid which contains 3 grains of Atoxyl on the first day, to repeat the injection on the fourth or fifth day, increasing the amount to $3\frac{1}{2}$ grains, and continuing the injection every third, fourth or fifth day, increasing each dose by $\frac{1}{2}$ grain until as much as 12 grains at a single dose is reached. Treatment of trypanosomiasis is continued for 2 or 3 months, after which an interval of 2 months is given, the series being then repeated. Atoxyl used in small doses, such as a grain twice a week to begin with and gradually increasing it, has been suggested in the treatment of leucæmia, pernicious anæmia, Hodgkin's disease — *The Hospital*, Sept 21 '07, p 657

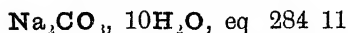
It has also been recommended (*B M J* '07, ii 685, 708) combined with Mercuric Chloride and Methylene Blue as a remedy for trypanosomiasis in veterinary work. A 1 p c aqueous solution of Methylene Blue prepared from a saturated Alcoholic solution is mixed with an equal quantity of a 1 in 500 aqueous solution of Mercuric Chloride. A dose of 10 c c is intravenously injected, and these intravenous injections have been repeated daily for 10 days.

Tests — Atoxyl dissolves readily in Water, forming a clear solution which possesses a neutral reaction towards Litmus paper, its aqueous solution reduces Potassium Permanganate Solution and Gold Chloride Solution. It yields with Ferrous Sulphate Solution a green precipitate, with Bromine Water a white precipitate, and with Sodium Hypobromite Solution a blackish-red coloration.

SOAMIN Sodium Para-aminophenylarsonate $\text{NaNH}_2\text{C}_6\text{H}_4\text{AsO}_3 \cdot 5\text{H}_2\text{O}$, eq 326.82 Is an organic combination of Arsenic, of which it contains 22.8 p c. It has been introduced for the treatment of syphilis and has been recently used with favourable results. It may be given by hypodermic injection, beginning with doses of 3 grains every third day and gradually raised to 10 grains every other day until a total of 100 grains has been given. As all the necessary physiological effects may be obtained with this dose it is not considered wise or necessary to push it any higher, the preparation should not be given by the mouth — *B M J* '08, ii 393

SODII CARBONAS.

SODIUM CARBONATE



FR, CARBONATI NEUTRI DI SODIUM CRISTALLISI OFFICINAR GER, NAIRIUM CARBONAT, ITAL, CARBONATO DI SODIO, SPAN, CARBONATO SODICO CRISTALLIZADO

Colourless, translucent, efflorescent, monoclinic crystals, possessing a somewhat caustic taste and an alkaline reaction. Sodium Carbonate (*BP*) contains 10 molecules of Water of crystallisation. The Sodium Carbonate official in the *USP* is monohydrated Sodium Carbonate containing 1 molecule of Water of crystallisation. The Sodium Carbonate (*PG*) is similar to the British and contains 10 molecules of Water of crystallisation.

It should be kept in well closed vessels, as it has a tendency to effloresce on exposure to dry air.

Solubility—5 in 8 of Water at 60° F, and measures 11, 12 in 1 of Water at 100° F, almost insoluble in Alcohol (90 p c).

Medicinal Properties—Antacid, but it is so apt to irritate that the Bicarbonate is almost invariably preferred. Externally, as a lotion (30 grains to 1 pint) in eczema.

Dose—5 to 30 grains = 0.32 to 2 grammes.

Prescribing Notes—*The Exsiccated salt may be given in the form of pills massed with 'Diluted Glucose'.*

143 grams of the crystallised salt are equal to nearly 53 grains of the Exsiccated salt.

20 of Sodium Carbonate are neutralised by 9.8 of Citric Acid, and by 10.5 of Tartaric Acid.

Official Preparation—Sodii Carbonas Exsiccatus used in the preparation of Extractum Ergotæ and many Sodium salts, also Liquor Magnesi Carbonatis and various Carbonates, etc. The Exsiccated Carbonate is used in the preparation of Pilula Ferri.

Not Official—Bileum Alkalinum, Baum Alevlin.

Foreign Pharmacopœias—Official in all.

U S has only Sodii Carbonas Monohydritus.

Tests—Sodium Carbonate when heated liquefies, loses its Water of crystallisation amounting to 62.93 p c and leaves a white anhydrous salt. The *USP* salt when heated to 100° C (212° F) loses its Water of crystallisation, equivalent to 14.52 p c. The *PG* states that 100 parts of the salt contain 37 p c of anhydrous Sodium Carbonate. On the addition of diluted Hydrochloric Acid it effervesces, giving off a colourless gas, which when passed into Lime Water yields a white precipitate soluble in a sufficient excess of the gas, and also soluble in a diluted mineral acid with effervescence. The resulting solution answers the tests distinctive of Sodium given under that heading. The *BP* requires that, with the exception of the Mercuric Chloride Solution test, it should answer the qualitative tests given under Sodii Bicarbonas. An aqueous solution is required to immediately yield a brownish-red precipitate on the addition of

Mercuric Chloride Solution It is presumably intended that the Magnesium Sulphate Solution test should also be excluded, for it may safely be taken that the words 'it should afford the reactions characteristic of Sodium and of Bicarbonates' apply to the qualitative tests for the latter. Sodium Carbonate dissolves readily in Water, yielding a clear solution which possesses a strongly alkaline reaction towards red Litmus paper. It is officially required to yield at least 98.01 p.c. of pure crystallised Sodium Carbonate, equivalent to 43.23 p.c. of pure anhydrous Sodium Carbonate as determined by titration with Volumetric Sulphuric Acid Solution. The mono-hydrated Sodium Carbonate of the *USP* is required to contain not less than 85 p.c. of pure anhydrous Sodium Carbonate, corresponding to not less than 99.5 p.c. of the crystallised mono-hydrated salt. The *PG* requires that it shall contain not less than 100.2 p.c. of pure crystallised Sodium Carbonate. The *BP* and *PG* methods of determination will be found in the small type below under the heading of Volumetric Determination. It will be observed that neither the *BP* nor the *PG* mentions a suitable indicator of neutrality. The *USP* requires that Methyl Orange Solution shall be used.

The more generally occurring impurities are such as are also found in the Bicarbonate, it may also contain Arsenic as an impurity. The tests given for the detection of the impurities under Sodium Bicarbonate may also be employed here, the modified Gutzzeit's test may also be used for the detection of Arsenic. Standards have been suggested (*CD* '08, 1796) of 10 parts per million for Lead and 2 parts per million for Arsenic. A standard solution for Chloride is 0.1 p.c., calculated as Sodium Chloride.

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by Hydrogen Sulphide T.S. either before or after acidulation with Acetic Acid, *PG*.

Barium Nitrate—An aqueous solution (1-20) acidulated with Acetic Acid should not be affected by T.S. of Barium Nitrate, *PG*.

Silver Nitrate—An aqueous solution (1-20) with excess of Nitric Acid should not give more than a whitish opalescence within 10 minutes with T.S. of Silver Nitrate, *PG*.

Sodium Hydroxide—Warmed with Sodium Hydroxide T.S. the salt should not evolve Ammonia, *PG*.

Volumetric Determination—6.9 c.c. of Sulphuric Acid Volumetric Solution should be necessary to neutralise 1 gramme of the salt, *BP*. Not less than 7 c.c. of Normal Volumetric Solution of Hydrochloric Acid, *PG*.

Preparation

SODII CARBONAS EXSICCATUS. Na_2CO_3 , eq 105.31. EXSICCATED SODIUM CARBONATE. DRIED CARBONATE OF SODIUM—*BP* '85

A white, amorphous, odourless powder, obtained by heating crystallised Sodium Carbonate until it loses 63 p.c. of its weight.

Dried Sodium Carbonate is not official in the *USP*. The mono-hydrated salt being the only one adopted. The dried salt is official in the *PG*, but it does not correspond to the completely dehydrated salt of the *BP*, as it is prepared by drying coarsely-powdered Sodium Carbonate at a temperature not

exceeding 25°C (77°F) until completely effloresced, protecting it from the dust, then at a temperature from 40° to 50°C (104° to 122°F) until it has lost half of its original weight. The official salt is obtained by heating Sodium Carbonate (temperature not stated) until it loses nearly 63 p c of its original weight, as stated above.

53 grams are equal to nearly 143 grains of crystallised salt.

Dose—3 to 10 grains = 0.2 to 0.65 gramme.

Foreign Pharmacopœias—Official in Austri, Dan, Fr, Ger, Hung, Jap, Russ, Spai, Swed and Swiss.

Tests—Dried Sodium Carbonate answers the tests distinctive of Sodium and of Carbonates given under the heading of Sodium Carbonas. Beyond the facts that it is required by the *BP* to yield not more than traces of Water when strongly heated, and that it is directed to be prepared as above, no indication is given of the requisite quantity of anhydrous Sodium Carbonate which it should contain. No method of determination is given in the *BP*. The *PG* requires that on the titration of a weighed quantity of 1 gramme of the dried Sodium Carbonate with Normal Volumetric Hydrochloric Acid, not less than 14 c c should be required for neutralisation. It should be free from the impurities mentioned under Sodium Bicarbonate, and also under Sodium Carbonate. It has been recommended that a limit of Water should be allowed and definitely stated.

Not Official

BALNEUM ALKALINUM—Crystals of Sodium Carbonate, 8 or 10 oz to 60 gallons of Water.

Used in scaly skin diseases.

BAIN ALCALIN—Crystallised Sodium Carbonate of commerce, 250 grammes dissolved in 1000 c c of Water and added to a bath.—*Fr*.

SODII CHLORIDUM.

SODIUM CHLORIDE

NaCl, eq 58.07

Fr, CHLORURI DI SODII OFFICINAT, Ger, Natriumchlorid
Ital, Cloruro di Sodio, Span, Cloruro Sódico

White, cubical crystals, or a white, crystalline powder, possessing a strong, saline taste, and neutral reaction. It is prepared by purifying common salt.

Solubility—1 in $2\frac{1}{4}$ of Water, 1 in $2\frac{1}{4}$ of boiling Water, 1 in 200 of Alcohol (90 p c).

Medicinal Properties—In small doses, stimulant and tonic, in larger doses, purgative and emetic, in the form of enema, anthelmintic. It is an important article of diet. A pint or more of **Normal Saline Solution** is injected intravenously, subcutaneously, or into the rectum, according to urgency, in shock or collapse due to hæmorrhage, and in uræmia, eclampsia and cholera. Locally, as a

fomentation to sprains and bruises Salt water baths (1 lb to 4 gallons) are tonic and stimulant, especially in children, and are useful in chronic rheumatism and gout Nasal injection of a saturated solution is useful in ozæna A recent cold is greatly relieved by douching the nostrils and gargling the throat with a weak solution of Salt, gargling is also serviceable in tonsillitis and chronic throat catarrh In case of a leech being swallowed a strong solution of Salt should be drunk, it is also a valuable antidote in poisoning by Silver Nitrate

Its value as an article of diet is well known Soldiers are supplied with it our army, 0.5 oz daily, the French, 0.5, Prussian, 0.87, Russian, 1.86, for a long time the Russian soldiers had salt-money given, and it was only when scurvy attacked them that the money was sold the salt given instead

Irrigation of the urethra with hot salt in treatment of gonorrhœa —*B M J E* '01, 11 60

Saline transfusion for prevention of shock during prolonged operations — *B M J* '01, 11 1189

Intravenous injection of normal saline solution in a severe case of hæmatemesis, recovery — *B M J* '02, 1 770

Since it is shown conclusively that a liberal allowance of salt may intensify the ascites, a diet relatively poor in Chlorides must be considered a useful therapeutic measure in such cases Good effects have been witnessed (*Pr lxxxiii* 699) from a cure by dechlorination in cases of ascites due to tubercular peritonitis, in pleurisy with effusion, and phlegmasia alba dolens

In desperate cases of hæmorrhage the subcutaneous transfusion of saline fluid should be practised, as it can be done without disturbing the patient — *B M J* '05, 1 68

Many of the symptoms, such as rigors and sweating, which are occasionally seen after intravenous transfusion of a solution of Sodium Chloride, are due to the chemical composition of the fluid being incorrect — *J* '05, 1 847

Bearing on this statement, it appears (*B M J* '04, 11 1198) that the proper strength for normal saline solution is 0.9 p.c

In collapse following severe hæmorrhage, intravenous transfusions with physiological salt solution should be performed as soon as possible — *L* '05, 1 854 1 to 10 oz of normal saline injected at one spot, the fluid being allowed to

in the treatment of collapse following the great loss of fluid caused of infants — *Pr lxxiv* 508 A case of fatal poisoning caused by injecting 500 c.c of an almost saturated solution from a stock bottle instead of the usual 0.9 p.c solution — *L* '05, 11 176 Subcutaneous injection of 100 to 300 c.c of Atlantic sea water, reduced by dilution to isotonicity with the blood, every 3 or 4 days in tuberculosis — *M P* '05, 11 383 Half a pint of normal saline solution injected twice daily in the treatment of congenital hypertrophic stenosis of the pylorus — *L* '05, 11 503

The injection of saline fluids may afford much assistance in surgical shock, but this fluid is expelled into the tissues more quickly the more profound the degree of shock, and the effect is therefore temporary and of no use whatever except to gain time, which may allow the superficial vessels to be made to relax or to relax spontaneously In this way saline injections may make all the difference between losing and saving a patient's life — *L* '05, 11 578

Injection of normal saline is stated (*B M J E* '05, 11 20) to show a superiority over other plans of treatment in delirium tremens

In the treatment of puerperal eclampsia as an intercellular transfusion, the solution used contains 1 drim each of Sodium Chloride and Sodium Acetate to a pint of Water The solution is sterilised, and at 100° F (37.7° C) is run into the areolar tissue beneath the breast or after delivery into the lax abdominal wall — *B M J* '05, 11 1635, *L* '01, 1 1682, *B M J* '01, 1 510, 958, 1144, '03, 1023, '03, 11 1332, 1378, 1408, *T G* '01, 616, 623

General formulas for saline solutions — *Pr lxxvii* 486, *P J* '99, 11 141.

In pneumonia — *B M J* '00, 11 900

In diabetic coma — *B M J* '03, 1 544

Dose—10 to 60 grains = 0.65 to 4 grammes, as a tonic, as an emetic, $\frac{1}{2}$ to 1 oz = 14.2 to 28.4 grammes

Official Preparation—Used in the preparation of Acidum Hydrochloricum, Hydrargyri Perchloridum, Hydrargyri Subchloridum, Sodii Bicarbonas and Sodii Sulphas

Not Official—Normal Saline Solution, Pulvis Salinus Anticholericus and Nebula Sodii Chloridi Composita

Foreign Pharmacopœias—Official in Austri, Belg., Dan., Dutch, Fr., Ger., Ital., Jap., Mex., Port., Russ., Spain, Swed., Swiss and U.S.

Tests—Sodium Chloride when heated decrepitates and at a red heat fuses. It dissolves readily in Water, forming a clear solution which is neutral in reaction towards Litmus paper and which yields the tests distinctive of Sodium given under that heading. It also yields on the addition of Silver Nitrate Solution a white curdy precipitate insoluble in Nitric Acid, readily soluble in Ammonia Solution and reprecipitated on acidification with Nitric Acid. When mixed with Sulphuric Acid it evolves Hydrochloric Acid gas, which immediately reddens a piece of moistened blue Litmus paper. When heated with Manganese Dioxide and Sulphuric Acid it evolves a yellowish gas, which first reddens and then bleaches a piece of moistened blue Litmus paper, and which instantly liberates Iodine from Potassium Iodide Solution, recognisable by the blue colour which it produces with Mucilage of Starch. Neither the *BP* nor the *PG* includes a requisite percentage of pure Sodium Chloride nor a method for its quantitative determination. The *USP* requires that the salt when dried should contain not less than 99 p.c. of pure Sodium Chloride as volumetrically determined by the method given in small type below under the heading of Volumetric Determination.

The more generally occurring impurities are Calcium and Magnesium, Bromides, Iodides and Sulphates. The *BP* includes also Potassium as a likely impurity. Calcium and Magnesium, if present, may be detected by the test with Ammonium Oxalate and Sodium Phosphate. Bromides or Iodides, if present may be detected by the test with Chlorine Water given below under that heading. Sulphates by the test with Barium Nitrate Solution given under the heading of Barium Nitrate. The *PG* includes a separate test for Iodine with Potassium Ferrocyanide Solution, the *USP* tests for Arsenic, Copper, Lead, Iron and Zinc by means of the time limit test with Hydrogen Sulphide. Each of these tests is described below under the headings Potassium Ferrocyanide and Hydrogen Sulphide. Standards have been suggested (*CD* '08, 1796) of 10 parts per million for Lead, and 1 part per million for Arsenic.

Hydrogen Sulphide—An aqueous solution (1-20) of the salt should not be affected by T.S. of Hydrogen Sulphide, *PG* slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*

Barium Nitrate—An aqueous solution (1-20) of the salt should not be affected by T.S. of Barium Nitrate, *PG*

Sulphuric Acid—An aqueous solution (1-20) of the salt should not be affected by diluted Sulphuric Acid, *PG*

Ammonium Oxalate—An aqueous solution (1-20) after the addition of Ammonia T S should not be affected by T S of Ammonium Oxalate, *P G*

Sodium Phosphate—An aqueous solution (1-20) after the addition of Ammonia T S should not be affected by T S of Sodium Phosphate, *P G*

Potassium Ferrocyanide—20 c c of an aqueous solution (1-20) should not be rendered blue by 0.5 c c of T S of Potassium Ferrocyanide, *P G*

Chlorine Water—If 2 grammes of the finely powdered salt be digested for some hours with 25 c c of warm Alcohol and, after cooling, the undissolved salt be removed by filtration, the filtrate evaporated to dryness, and the residue dissolved in 5 c c of Water, and if 1 c c of Chloroform be added, and Chlorine Water which has been diluted with twice its volume of Water, cautiously introduced, drop by drop, with constant agitation, the Chloroform should not acquire a violet, yellow, or orange colour, *U S P*

Volumetric Determination—If 1 gramme of well dried Sodium Chloride be dissolved in sufficient Distilled Water to measure 100 c c, and 10 c c of the solution be mixed with a few drops of Potassium Chromate T S, it should require not less than 17 (17.05) c c of Tenth-normal Volumetric Silver Nitrate Solution to produce a permanent red colour, *U S P*

Not Official

NORMAL SALINE SOLUTION (also called Physiological Salt Solution)

—Sodium Chloride, 78.75 grains, boiled and cooled Water, to 20 fl oz (or in any case, sterilised)

On the authority of Professor Stirling, of Manchester, the percentage of Sodium Chloride in human blood is taken to be 0.9 p c. The usual figure 0.6 p c is for cold-blooded animals, and was calculated on the blood of a frog.

Solutions the osmotic pressure of which is the same as that of blood plasma, are said to be isotonic with the blood. If a solution the osmotic pressure of which is markedly less than that of blood plasma be used, many of the red corpuscles may be destroyed. The salt in greatest amount in the plasma is Sodium Chloride, and therefore in making isotonic solutions this salt is generally used. A solution containing 0.9 p c Sodium Chloride gives the same osmotic pressure as plasma, and is therefore isotonic with the blood.

PULVIS SALINUS ANTICHOLOERICUS (*Stevens*)—Sodium Bicarbonate, 30 grains, Sodium Chloride, 20 grains, Potassium Chlorate, 7 grains, for 1 dose.

Given frequently in a small tumbler of Water in diarrhoea and cholera.

NEBULA SODII CHLORIDI COMPOSITA—Sodium Chloride, 1, Sodium Bicarbonate, 1, Borax, 1.

1 teaspoonful to be dissolved in a pint of warm Water, and used as a spraying solution.—*University*

Sodium Bicarbonate, 1.50, Sodium Chloride, 0.75, Borax, 1.50, Distilled Water, warm, *q s* to produce 100.—*B P C*

Not Official

SODII CINNAMAS

SODIUM CINNAMATE

$\text{Na C}_6\text{H}_5\text{O}_2$, eq 168.83

A white, granular, amorphous powder, having a faint cinnamon-like odour, and a faintly alkaline reaction. Soluble 1 in 11 of Water, 1 in 160 of Alcohol (90 p c). It has been used in phthisis and in cancer, as an intramuscular or intravenous injection (15 to 30 minims of a 10 or 20 p c aqueous or Glycerin solution). In ophthalmic surgery it has been employed in the form of a 1 p c (7 to 8 minims) aqueous solution by subconjunctival injection.—*L* '02, 11 66, 67, 1755, *B M J E* '01, 1 67, '02, 1 28, *P J* '02, 1 350, *C D* '02, 11 155.

In tuberculosis, commencing with doses of 1 mg = $\frac{1}{10}$ cc of 1 p c solution, increasing the dose by $\frac{1}{2}$ to 1 mg until 10, 15, or even 20 mg are reached, the injections being given 3 times a week —*B M J E* '04, 1 71

Though not a specific, it is a useful adjunct to treatment of tuberculosis. Dose, 20 to 50 mg (maximum) Leucocytosis produced by about 20 mg —*I*, '04, 11 1136, *B M J E* '05, 1 24

In the treatment of cancer it is administered once or twice a week in the form of a 10 p c Glycerin solution in doses of 30 minims (1.8 cc) hypodermically. The hypodermic use of a 22 p c aqueous solution of Sodium Ortho coumarate in doses of 25 minims (1.5 cc), administered at the same interval, is also referred to —*B M J* '05, 1 927

In a plea for more extended use of treatment by hypodermic injection a reference is made (*L* '05, 1 1340) to the successful employment of this salt by Landerer in 1893 in the treatment of pulmonary tuberculosis.

A further reference to the use of these salts in cancer, and to a serum prepared from tuberculous horses after repeated injections with Sodium Cinamate, is in *L* '05, 11 393

An interesting point concerning the solubility of this salt and the preparation of the solution is dealt with in a paper on Solubility appearing in the *P* / [4], 20, 786 *C D* '05, 11 783

The most active chemical substance examined was Cinnamic Aldehyde, its administration is, however, exceedingly painful, and had to be temporarily abandoned.

Although the correct figure for the solubility of this salt is 1 in 11, a clear solution may be prepared by the aid of heat of a strength of 1 in 10, but it requires very careful handling, and the friction of the stopper of the bottle is very often sufficient to cause the salt to crystallise out. For hypodermic use the solution should be a little weaker than 10 p c. Some authorities have claimed that a clear permanent 10 p c solution may be prepared in absolute Glycerin, but this is contrary to our experience. The solution, though clear when first prepared, develops crystals in the course of a few days, and the specimens before us now are practically a solid mass of crystals. The salt has assumed importance owing to its hypodermic employment in pulmonary tuberculosis and cancer. Having had a large number of these solutions to prepare, this experience of the behaviour of the salt may prove useful to others.

Dose —2 to 5 grains = 0.13 to 0.32 gramme

It was introduced commercially as **Hetol**

Tests—Sodium Cinamate when heated yields an aromatic odour somewhat resembling Bitter Almonds, and when ignited with free access of air leaves a more or less blackish residue which when dissolved in Water possesses a strong alkaline reaction towards red Litmus paper, and which effervesces on the addition of a diluted mineral acid.

The salt dissolves in Water, yielding a clear solution which is neutral in reaction towards Litmus paper. An aqueous solution yields with Ferric Chloride T.S. a yellow precipitate, and a white precipitate with Manganese Chloride Solution, which soon becomes crystalline. If the aqueous solution be sufficiently concentrated it yields on acidification with Diluted Sulphuric Acid a white crystalline precipitate soluble in Ether. If the ethereal solution be separated, washed till free from mineral acid and evaporated spontaneously, it yields a crystalline residue which should possess a m.p. of about 132° C (269.6° F), the aqueous portion remaining after the removal of the precipitated acid will answer the tests distinctive of Sodium given under that heading. When oxidised with Potassium Permanganate it yields an odour of Benzaldehyde. The amount of pure Sodium Cinamate contained in a specimen may be quantitatively determined by titration with Tenth normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as an indicator of neutrality, sufficient Ether should be added to hold the liberated Cinnamic Acid in solution during the titration, and to prevent it masking the end reaction. 1 cc of Tenth normal Volumetric Sulphuric Acid Solution corresponds to 0.016883 gramme of pure anhydrous Sodium Cinamate. Free Cinnamic Acid, if present, may be determined, previous to the above volumetric determination, by titrating with Tenth

normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. Each c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution used $\frac{1}{10}$ gramme of absolute Cinnamic Acid. In the event of free Cinnamic Acid being present, the number of c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution required to neutralise this free Cinnamic Acid must be deducted from the total number of Tenth-normal Volumetric Sulphuric Acid Solution required to complete the titration with Methyl Orange Solution before the result is calculated into terms of Sodium Cinnamate. The aqueous solution when acidified with diluted mineral acids, though affording a white crystalline precipitate, should yield no effervescence, indicating the absence of Carbonates. If the aqueous solution be acidified with diluted Nitric Acid, and the liberated Cinnamic Acid be separated by filtration, the filtrate should yield at the most but a slight turbidity with either Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of Chlorides and Sulphates.

Hetol-Caffeine (Caffeine Sodium Cinnamate)—An amorphous, bitter powder, **Heto-Cresol** (Meta-cresolic Ester of Cinnamic Acid) and **Hetoform** (Bismuth Cinnamate) are compounds which have received notice in medical literature.

INJECTIO SODII CINNAMATIS—A sterilised 10 p.c. aqueous solution of Sodium Cinnamate.

Dose—16 minims = 1 c.c., hypodermically.

Glycerin has been recommended as a solvent for Sodium Cinnamate. The solution can be readily sterilised, but a 10 p.c. solution, though bright when first made, soon crystallises out.

This suggestion has been incorporated in the *B.P.C.* in the form of **Glycerinum Sodii Cinnamatis**, Sodium Cinnamate, 5, Glycerin, 95.

Not Official SODII CITRAS

SODIUM CITRATE

$\text{Na}_3\text{C}_6\text{H}_5\text{O}_7, 5\frac{1}{2}\text{H}_2\text{O}$, eq. 354.60

A white, granular powder, possessing a cool, saline taste and a faint, caramel-like odour.

It should be kept in well-closed bottles, as it has a tendency to slowly effloresce on exposure to dry air. It dissolves 5 in 9 of Water, insoluble in Alcohol (90 p.c.), and in Ether.

Tests—Sodium Citrate when heated loses its Water of crystallisation, when heated to dull redness it is decomposed, and on ignition leaves a carbonaceous residue which, when dissolved in Water, possesses a strongly alkaline reaction. This residue effervesces on the addition of diluted Hydrochloric Acid, and yields a solution giving the tests distinctive of Sodium mentioned under that heading. The salt dissolves readily and completely in Water, forming a clear solution slightly alkaline in reaction towards red Litmus paper, and which should not be coloured red by a drop of Phenolphthalein Solution. The aqueous solution affords upon boiling with Calcium Chloride Solution a white granular precipitate, insoluble in Potassium Hydroxide but soluble in Ammonium Chloride Solution. The *U.S.P.* requires the salt to contain not less than 97 p.c. of pure crystallised Sodium Citrate as volumetrically determined by the titration of the solution of the alkaline residue left on ignition. A weighed quantity of the salt is thoroughly charred at a dull red heat, the residue extracted with boiling Water till the washings fail to react with Methyl Orange T.S., and the mixed filtrate and washings are titrated with Semi-normal Volumetric Sulphuric Acid Solution, employing the above indicator to ascertain the point of neutrality, not less than 16.4 c.c. of the Semi-normal Volumetric Sulphuric Acid Solution should be necessary.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron, Zinc and Carbonates. Arsenic, if present, may be detected by the modified Gutzzeit's test. A 5 p.c. w/v aqueous solution of the salt should neither yield a distinct coloration nor turbidity on the addition of Hydrogen Sulphide to the solution acidified with diluted Hydrochloric Acid, nor a decided coloration or turbidity when the solution is subsequently made alkaline with Ammonia, indicating the absence of Copper, Lead, Iron and Zinc. A 1 in 20 aqueous solution of the salt should not yield an effervescence on the addition of a mineral acid, indicating the absence of Carbonates. The Phenolphthalein test given above also affords an indication of Carbonates, if present.

Its addition in the proportion of 1 grain to the oz., or, if necessary, 1½ to 2 grains, is found (*P* lxiv 221) to render the curd of Milk more easily digestible, but the salt tends to produce constipation.

The addition of 1 grain to 1 oz. Milk, increased if necessary to 2 or 3 grains to the oz., renders the curd of Milk more easily digested — *L* '05, ii 364. In the feeding of the infant, 1 grain to the oz. of Milk is prescribed in a teaspoonful of Water and added by the mother to the bottle before every meal — *B M J* '05, ii 1022.

While not denying the empeptic action which has been ascribed to this drug in infantile gastro enteritis, its chief value (*L* '05, ii 192) is its power of acting as an anti emetic in the digestive troubles of bottle fed infants, or in those who, although suckled, vomit for no definite reason.

Good results in dyspepsia — *L* '07, i 309.

Official in U S

SODIUM COUMARATE There are three isomeric Coumaric acids (ortho, meta, and para), forming salts with a Sodium base, known as Sodium Ortho Coumarate, Sodium Meta coumarate, and Sodium Para coumarate. The employment of the Sodium salt of Cinnamic Acid in the treatment of cancer (*L* '02, ii 66 *B M J* '05, i 927) has led to a search for a similar substance possessing an increased physiological action. This has been found (*B M J* '05, i 1143) in Coumaric Acid, a substance having the structure of Cinnamic Acid with a hydroxyl substituent. A 22 p.c. aqueous solution of the **Sodium Ortho-coumarate** containing a slight excess of the free Ortho coumaric Acid was tried. The solution induced a rapid physiological action, the leucocytosis being well marked and resembling that effected by the Cinnamate. An 8 p.c. solution of the less soluble **Sodium Para-coumarate** was employed, the results tending to show the action was of a similar nature, but rather less intense than that produced by the ortho salt.

A 20 p.c. aqueous solution of the **Sodium Meta-coumarate** was used, and showed a very marked physiological action, being apparently even more active than the ortho compound. The three acids are certainly physiologically active, but it must be left for further experiments to decide which of these is likely to prove to be the most serviceable therapeutic agent.

Beneficial in cancer, even in cases of worst possible type — *L* '07, ii 690.

SODII ETHYLATIS LIQUOR.

SOLUTION OF SODIUM ETHYLATE

A pale yellow, viscid, alcoholic liquid, prepared by dissolving 22 grains of clean, bright, metallic Sodium in 1 fl. oz. of Absolute Alcohol, care being taken to keep the contents of the flask cool during the reaction. It is officially described as a colourless liquid, but even when freshly prepared it scarcely answers this description, being usually of a pale straw-colour, and becoming yellowish-brown on keeping, and when traces of aldehyde are present in the Alcohol.

the change of colour is more rapid and occurs to a much greater extent, producing a deep brown

This solution should be recently prepared, and should be preserved in well-stoppered bottles of a dark amber tint. It contains 18 p.c. of the solid substance, C_2H_3ONa .

If the Sodium be not bright, it is advisable to wash it with a little Absolute Alcohol before commencing to make the Liquor.

Medicinal Properties—Caustic, used in the treatment of naevus, nasal polypus, ozoena, warts and lupus—*L* '78, ii 625, '81, i 168, 242, *B M J* '85, ii 344, '88, ii 762

Successful in multiple circumscribed lymphoma—*L* '07, i 943

It may be applied by means of a glass rod, camel's-hair brush, or a quill pen. Tincture of Opium may be added to relieve the pain, but not Chloroform, as it makes an explosive mixture.

Tests—Sodium Ethylate Solution is required by the *BP* to possess the sp. gr. of 0.867. It boils when heated, emitting vapours possessing an alcoholic odour, a white residue remaining, which undergoes hydrolysis when heated with Water, yielding a solution which possesses a strongly alkaline reaction towards red Litmus paper and which produces a strong red coloration with Phenolphthalein Solution. If a portion of this solution be evaporated to dryness it leaves a residue which should answer the tests distinctive of Sodium Hydroxide given under the heading of Soda Caustica. The white residue left on the evaporation of the Alcohol chars when strongly heated.

SODII HYPOPHOSPHIS.

SODIUM HYPOPHOSPHITE

$NaPH_2O_2$, eq 87.44

FR, HYPOPHOSPHITE DE SODIUM, GER, NATRIUMHYPOPHOSPHIT,
ITAL, IPOFOSFITO DI SODIO, SPAN, HIPOFOSFITO SODICO

Colourless, translucent, deliquescent, prismatic crystals, or as a white, granular powder, ' a slightly bitterish, saline taste. It is obtained by the interaction of Sodium Carbonate and Calcium Hypophosphite.

Sodium Hypophosphite, when mixed with an equal quantity of Sodium Nitrate, forms a highly explosive mixture—*Y B P* '87, 21.

It should be kept in well-closed vessels in a cool atmosphere and protected as far as possible from contact with the air, as it is stated to be of a deliquescent nature. The crystals or powder deliquesce slowly in very hot weather, but as soon as it cools [say to 18–3° C (65° F)] the salt dries up again. It should be handled with caution, as it is readily oxidised, and when brought into contact with powerful oxidising agents the temperature rises so rapidly that an explosion is liable to result. The formula given in the *BP* shows the official salt to be anhydrous, the *USP* formula is given with 1 molecule of Water of crystallisation, the salt is not official in the *PG*.

Solubility.—1 in 1 of Water, 1 in 2 of Glycerin, almost entirely 1 in 20 of Alcohol (p.c.).

Medicinal Properties—Similar to those of Calcium Hypophosphite

Dose—3 to 10 grains = 0.2 to 0.65 gramme

Not Official—Syrupus Sodii Hypophosphitis

Foreign Pharmacopœias—Official in Belg., Dutch, Fr., Ital., Mex., Port., Span. and U.S.

Tests—Sodium Hypophosphite when heated evolves spontaneously inflammable Hydrogen Phosphide gas and Hydrogen, the *U.S.P.* states that when heated in a test-tube the salt first loses its Water of crystallisation, and at about 200° C (392° F) it is decomposed, evolving Hydrogen and Hydrogen Phosphide gas, which burns spontaneously with a bright yellow flame. The salt answers the tests distinctive of Sodium given under that heading. It dissolves readily in Water, forming a clear solution which is neutral or only faintly alkaline to Litmus paper. The solution yields with warm Copper Sulphate Solution a reddish brown precipitate of Cuprous Hydride, and on boiling evolves Hydrogen. It rapidly decolorises solution of Potassium Permanganate. The diluted aqueous solution acidulated with Diluted Sulphuric Acid yields on the addition of Silver Nitrate Solution a white precipitate rapidly turning from brown to black, owing to its reduction to metallic Silver. On the addition of Mercuric Chloride Solution to a 5 p.c. aqueous solution of the salt acidulated with Hydrochloric Acid a white precipitate is produced, changing rapidly to grey, owing to its reduction to metallic Mercury. The *B.P.* utilises its reducing action on Potassium Permanganate Solution as a basis for a method of determination, requiring that when a weighed quantity of 0.5 of a gramme of the salt is boiled for 10 minutes with a solution of 1.15 grammes of Potassium Permanganate in 25 c.c. of Water and filtered, a nearly colourless filtrate should be yielded. No statement is made respecting the amount of pure Sodium Hypophosphite which compliance with this test indicates. It has been recommended that this Permanganate test should be replaced by one based on the work of Jowett, the method is described under Calcium Hypophosphite. The *U.S.P.* states that the salt should contain not less than 98 p.c. of pure crystallised Sodium Hypophosphite, but gives no method for its determination.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Zinc, Calcium Magnesium, alkali or alkali Carbonate, Chlorides and Sulphates, Phosphates and Phosphites. Arsenic, if present, may be detected by the modified Gutzeit's test given in small type below under that heading, Copper, Lead, Iron and Zinc, if present, by the Hydrogen Sulphide test. The aqueous solution of the salt should afford no distinct turbidity with Ammonium Oxalate Solution after the addition of a little Ammonium Chloride Solution. If the mixture be allowed to stand for some time and filtered it should yield little or no turbidity with Sodium Phosphate Solution, indicating the absence of Calcium and Magnesium. The aqueous solution should neither be coloured red on the addition of Phenolphthalein T.S. nor should it effervesce on the addition of a

diluted mineral acid. When acidified with Diluted Nitric Acid, the aqueous solution should afford no pronounced turbidity or precipitate with either Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. It is officially required to yield no precipitate with Lead Acetate Solution, indicating a limit of Phosphates and Phosphites. The remarks on the Lead Acetate test will be found under Calcium Hypophosphorus.

Hydrogen Sulphide—An aqueous solution (1-20) acidulated with Hydrochloric Acid should not respond to the time-limit test for heavy metals, *U S P*.

Gutzeit's Test—If 5 c.c. of an aqueous solution of the salt (1-10) be measured into a beaker containing 3 c.c. of Nitric Acid diluted with about 10 c.c. of Water and evaporated to dryness on a water-bath, the residue should not respond to the modified Gutzeit's test for Arsenic, *U S P*.

Not Official

SYRUPUS SODII HYPOPHOSPHITIS—Dissolve 160 grains of Sodium Hypophosphite in 3 fl. drms. of Distilled Water, filter, and wash the filter with Distilled Water 1 fl. drm. To the filtered solution add sufficient Syrup to produce 20 fl. oz. Each fl. drm. contains 1 grain of Sodium Hypophosphite.—*B P C Formulary* 1901.

Dose—1 to 4 fl. drms. = 3.6 to 14.2 c.c.

Official in Mex.

Dissolve 2 of Sodium Hypophosphite in 2 of Distilled Water and add sufficient Syrup to the filtered solution to make 100.—*B P C*.

SODII IODIDUM.

SODIUM IODIDE

NaI, eq 148.78

FR, IODURE DE SODIUM, GER, NATRIUMJODID, ITAL, JODURO DI SODIO, SPAN, YODURO SODICO

Colourless, cubical crystals, or an odourless, white, crystalline, or fine powder, possessing a somewhat bitter, saline taste. It is stable in moist air, becoming partially decomposed, it should therefore be kept in well-closed bottles and in a cool place. It is officially described as a 'dry' powder, but commercial samples vary much in the proportion of Water which they contain, from 10 to 20 p.c.

Solubility.—11 in 6 of Water, and measures 10½, 1 in 3 of Alcohol (90 p.c.), 1 in 1 of Glycerin.

Medicinal Properties—Given in the same doses as, and for purposes similar to those of, Potassium Iodide, is more readily tolerated by the stomach, and is less depressant.

5 to 10-grain doses for long continued administration, combined in the earlier stages with Ammonia, and in the later with from 3 to 5 minims of Fowler's Solution, in the treatment of pain at the heart after influenza.—*L* '99, 11 1081.

Dose.—5 to 20 grains = 0.32 to 1.3 gramme

Foreign Pharmacopœias—Official in all except Port

Tests—Sodium Iodide melts when strongly heated. It dissolves readily in Water, forming a clear solution which is faintly alkaline to red Litmus paper. It answers the tests distinctive of Sodium given under that heading. The aqueous solution yields with Silver Nitrate Solution a yellow curdy precipitate insoluble in Nitric Acid, practically insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution. With Mercuric Chloride T.S. it yields a scarlet precipitate soluble in excess of the reagent and very soluble in excess of Sodium Iodide. With Lead Acetate Solution it yields a yellow precipitate soluble in diluted Nitric Acid, and also in boiling Water, from which solution on cooling it recrystallises in beautiful crystalline scales. When the aqueous solution is mixed with Chlorine Water it yields a reddish brown coloration, and on shaking the liquid with Carbon Bisulphide the latter solution is coloured a deep violet tint. The dried salt is required to contain not less than 98.9 p.c. of pure Sodium Iodide as determined volumetrically by titration with Tenth normal Volumetric Silver Nitrate Solution, *see* below under the heading Volumetric Determination. It is officially required to lose not more than 5 p.c. of Water when dried at 120° C (248° F). Commercial samples vary much in the proportion of Water which they contain. The *U.S.P.* requires that the salt shall contain at least 98 p.c. of pure Sodium Iodide as volumetrically determined by the method given in small type below under the heading of Volumetric Determination. It does not state a limit of loss of weight when dried. The *P.G.* requires that 100 parts shall contain at least 95 parts of anhydrous salt, the *P.G.* does not include a limit of moisture.

The more generally occurring impurities are Arsenic, Copper, Lead and Iron, Ammonium, Barium, Magnesium and Potassium, Bromates and Carbonates, Cyanides, free Iodine, Iodates, Chlorides, Bromides, Thiosulphates and Sulphates. Arsenic, Copper, Lead and Iron, if present, may be detected by the time-limit test for heavy metals mentioned in small type under the heading of Hydrogen Sulphide. The salt should not evolve an odour of Ammonia when boiled with Potassium Hydroxide Solution, nor should the issuing vapour have an alkaline reaction upon moistened red Litmus paper, indicating the absence of Ammonium salts. The aqueous solution of the salt should yield no reaction with the test for Barium described below under the heading of Potassium Sulphate. It should yield a scarcely perceptible turbidity on the addition of Ammonium Oxalate Solution, and if the mixture be set aside for some time and filtered, the filtrate should yield little or no turbidity on the addition of Sodium Phosphate Solution, indicating the absence of Calcium and Magnesium. It should afford no reaction for Potassium when examined by the test given below under the heading of Sodium Bitartrate. The *B.P.* includes a test for Bromates, presumably that mentioned in the Appendix is to be applied to this salt. Its aqueous solution should not effervesce on the addition of a diluted mineral acid, indicating the absence of Carbonate. It should yield no indication of the presence of Cyanide when examined by the test with Ferrous Sulphate, Ferric Chloride and Alkali Hydroxide described

below No indication of free Iodine should be yielded when $\frac{1}{2}$ gramme of the salt is dissolved in 10 c c of Distilled Water, which has been previously boiled and cooled in a small flask, as indicated by the solution not acquiring a yellow tint, nor should the solution acquire a yellow colour within half a minute after an addition of 2 drops of Diluted Sulphuric Acid, indicating the absence of Iodate Chlorides, Bromides and Thiosulphates, if present, may be detected by the test with Tenth-normal Volumetric Silver Nitrate Solution and Ammonia described below in small type An aqueous solution of the salt should not afford a distinct turbidity with Barium Chloride Solution, indicating the absence of more than traces of Sulphates The *USP* includes a test for limit of alkali, requiring that a drop of Phenolphthalein T S added to a solution of 1 gramme of the salt in Water, and containing 0.1 c c of Tenth-normal Volumetric Sulphuric Acid Solution should not produce a red coloration The *PG* and the *USP* both include a test for Nitrates and Nitrites, that of the former Pharmacopœia fixing a limit for these impurities by the adoption of a time limit during which no coloration should become apparent, the tests are described in small type below under the headings of Aluminium Wire and Potassium Hydroxide, or Zinc filings, powdered Iron and Sodium Hydroxide, the *USP* employing the former mixture, the *PG* the latter The *PG* includes a separate test for Iodine described below under the heading of Potassium Ferrocyanide Solution The alternative method for the determination of Potassium Iodide in a mixture of Potassium Chloride, Bromide and Iodide described under the heading of Potassium Iodide is equally applicable in the case of the Sodium Iodide

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by T S of Hydrogen Sulphide, *PG*, slightly acidulated with Hydrochloric Acid it should not respond to the time-limit test for heavy metals, *USP*

Barium Nitrate—An aqueous solution of the salt (1-20) should not be affected by T S of Barium Nitrate, *PG*

Ferrous Sulphate, Ferric Chloride, and Alkali Hydroxide—If an aqueous solution of the salt (1-20) be gently warmed with a crystal of Ferrous Sulphate, 1 drop of Ferric Chloride T S and Sodium Hydroxide T S the mixture should not be coloured blue on supersaturating with Hydrochloric Acid, *PG*, 5 c c of the aqueous solution, gently heated with 1 drop each of Ferrous Sulphate and Ferric Chloride T S and 0.5 c c of Potassium Hydroxide T S, should not develop a blue colour after acidulating with Hydrochloric Acid, *USP*

Sulphuric Acid—If Starch T S and diluted Sulphuric Acid be added as quickly as possible to a freshly prepared solution (1-10) of the salt in previously boiled and cooled Water, the solution should not be immediately coloured, *PG* The *USP* states that the solution of 0.5 gramme of the salt in 10 c c of previously boiled and cooled Distilled Water should not have a distinct yellow tint, nor should it acquire a yellow colour within half a minute after the addition of 2 drops of Diluted Sulphuric Acid (which should be free from Sulphurous Acid or Nitrous Acid)

Phenolphthalein—A solution of 1 gramme of the salt in Water with 0.1 c c of Tenth-normal Sulphuric Acid Volumetric Solution added should give no red colour with a drop of Phenolphthalein T S even after heating, *USP*

Sodium Bitartrate—A solution of 1 gramme of the salt in 1 c c of Water should yield no precipitate with 1 c c of Sodium Bitartrate, *USP*.

Potassium Sulphate—10 c.c. of an aqueous solution (1-20) acidulated with Hydrochloric Acid should not be rendered turbid by the addition of 1 c.c. of T S of Potassium Sulphate, *U S P*

Potassium Ferrocyanide—20 c.c. of an aqueous solution (1-20) after the addition of a few drops of Hydrochloric Acid should not be rendered blue by 0.5 c.c. of Potassium Ferrocyanide, *P G*

Zinc Filings, Powdered Iron, and Sodium Hydroxide—1 gramme of the salt warmed with 5 c.c. of Sodium Hydroxide T S and a mixture of 0.5 gramme of Zinc filings and powdered Iron, should not evolve Ammonia, *P G*

Aluminium Wire and Potassium Hydroxide—If to 1 gramme of the salt contained in a test tube of about 40 c.c. capacity, 5 c.c. of Water, 5 c.c. of Potassium Hydroxide T S, and about 0.2 gramme of Aluminium wire be added, and if in the upper portion of the test tube a pledget of purified Cotton be inserted, and over the mouth there be placed a piece of moistened red Litmus paper, then if the tube be heated upon a water bath for 15 minutes, no blue coloration of the paper should be discernible, *U S P*

Tenth-normal Volumetric Silver Nitrate and Ammonia—If 0.2 gramme of the salt (dried, *P G*) be dissolved in 2 c.c. of Ammonia T S and 15 c.c. (14 c.c., *P G*) of Tenth normal Silver Nitrate Volumetric Solution be added, then after thoroughly agitating and filtering, the filtrate upon supersaturating with Nitric Acid, should not become more than slightly turbid, nor should any darkening appear within 10 minutes, *P G* and *U S P*

Volumetric Determination—A solution of 1 gramme of the salt [dried at 120° C (248° F)] in Water should require the addition of not less than 66.5 c.c. of Volumetric Solution of Silver Nitrate for complete precipitation, *B P*, a solution of 0.5 gramme of the well dried salt in 10 c.c. of Water with about 5 drops of Potassium Chromate T S added, should require not more than 34.6 c.c. nor less than 33 c.c. of Tenth normal Silver Nitrate Volumetric Solution to produce a permanent red colour (corresponding to at least 98 p.c. of pure Sodium Iodide), *U S P*

Not Official

Rubidium Iodide has been used for similar purposes to the Potassium Iodide

Dose—5 to 20 grains = 0.32 to 1.3 grammes

SODII NITRIS.

SODIUM NITRITE

NaNO2, eq 68.58

White, or yellowish-white, fused pencils or sticks, with a crystalline fracture, or a whitish, deliquescent crystalline powder, possessing a mild, saline taste, and an alkaline reaction

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the air and light, it has a tendency to deliquesce and to become gradually oxidised. It is frequently found in commerce fused into sticks, with a crystalline fracture. It is prepared by fusing Sodium Nitrate with reducing substances, such as metallic Lead, Barium Sulphide, etc., but if the reduction is carried too far, free alkali is formed and afterwards becomes carbonated

Solubility—5 in 6 of Water, 1 in 50 of Alcohol (90 p.c.)

Medicinal Properties—Vaso-dilator and antispasmodic. Used with the object of warding off the attack in *angina pectoris* and *asthma*, as well as relieving the symptoms during an attack, also

in migraine and hemicrania if accompanied by facial pallor. It is not so rapid in its action as Amyl Nitrite, but is more persistent and more gentle. It is of great service in lowering arterial tension in renal cirrhosis.

Closely approaches the action of Nitroglycerin, but without its objectionable features.—*P.* xxx 179

1 to 4-grain doses every 3 or 4 hours combined with Aromatic Spirit of Ammonia, and sometimes with $\frac{1}{16}$ grain of Morphine Hydrochloride in angina pectoris.—*P.* li 348

Dose.—1 to 2 grains = 0.06 to 0.13 gramme

Official Preparation.—Used in the preparation of Liquor Ethyl Nitritus

Antidotes.—Emetics, fresh air, recumbent position, Ergot, and Atropine

Foreign Pharmacopœias.—Official in Swiss and U.S.

Tests.—Sodium Nitrite when heated melts, and at red heat is decomposed. It dissolves readily in Water, forming a clear solution, slightly alkaline in reaction towards red Litmus paper. It answers the tests distinctive of Sodium given under that heading. Potassium Iodide Solution and Starch Mucilage when added to an aqueous solution yield on the addition of a few drops of Diluted Sulphuric Acid a blue coloration. With Ferrous Sulphate Solution and Acetic Acid the aqueous solution affords a deep brown colour, the salt evolves red fumes when mixed with Diluted Sulphuric Acid. It is officially required to contain not less than 95 p.c. of Sodium Nitrite as gasometrically determined by measuring the quantity of Nitrous Acid gas evolved, when a solution of a weighed quantity is introduced into a nitrometer containing a saturated solution of Bromine, Potassium Iodide and Diluted Sulphuric Acid, a weighed quantity of 0.1 of a gramme of the salt should liberate at the ordinary temperature 15–5° C (60° F) and pressure (30 in. or 760 mm. of Mercury) not less than 32.5 c.c. of gas, which should be almost completely absorbed by concentrated Ferrous Sulphate Solution. It has been suggested that the Volumetric Permanganate test is better than the gasometric method now official. This is a very appropriate recommendation, the volumetric test being readily and quickly carried out and being much better adapted to the purpose than the obsolete method of gasometric analysis. Good commercial samples commonly yield 98 p.c. of pure Sodium Nitrite. The 17th Edition of *Squire's Comparison* stated that in the absence of a nitrometer it may be readily estimated with a standard solution of Potassium Permanganate, 0.1 of a gramme of pure Sodium Nitrite being equal to 29 c.c. of Tenth-normal Volumetric Potassium Permanganate Solution (containing 3.156 grammes of Potassium Permanganate in 1 litre) or to 9.1 c.c. of Liquor Potassii Permanganatis, *B.P.* The *U.S.P.* requires the salt to contain not less than 90 p.c. of pure Sodium Nitrite as volumetrically determined by the addition of an excess of Tenth-normal Volumetric Potassium Permanganate Solution, the excess of the latter with Tenth-normal Volumetric Oxalic Acid Solution as described under the heading of Volumetric Determination in small type below.

The more generally occurring impurity is Lead. The *B.P.* requires

that on the addition of Diluted Sulphuric Acid to the aqueous solution not more than the slightest traces of a precipitate should be produced, indicating the absence of Lead, the *U S P* employs the time limit test with Hydrogen Sulphide for the detection of heavy metals, such as Arsenic, Copper, Lead, Iron and Zinc

Time-limit Test—If 1 gramme of the salt be dissolved in 20 c.c. of Diluted Hydrochloric Acid, and heated sufficiently to expel the gases, the resulting solution after restoring it to its original volume should not respond to the time limit test for heavy metals, *U S P*

Volumetric Determination—If to 30 c.c. of Tenth normal Volumetric Potassium Permanganate Solution, diluted with about 150 c.c. of Distilled Water, 5 c.c. of Sulphuric Acid and 10 c.c. of a solution of 1 gramme of Sodium Nitrite in sufficient Distilled Water to make 100 c.c. be successively added, the liquid brought to a temperature of 40° C (104° F) and allowed to stand for 5 minutes, not more than 3.75 c.c. of Tenth normal Oxalic Acid Volumetric Solution should be required to decolorise the solution (each c.c. of Tenth normal Potassium Permanganate consumed corresponding to 0.004285 gramme of pure Sodium Nitrite, *U S P*)

Not Official

SODIUM PERBORATE—A white, crystalline substance which possesses the property of evolving Hydrogen Peroxide when dissolved in cold Water.—*B M J* '05, 1 310, *P J* '05, 1 193

SODIUM PHENOL SULPHO-RICINATE—This has been used as a spray in the treatment of pyiloinitis.—*B M J* '04, 11 1225

SODIUM PHENYLPROPIOLATE—Phenyl propionic Acid is prepared by treating Methyl dibromocinnamate with alcoholic solution of Potassium Hydroxide. The pure acid has a m.p. 136° to 137° C (276.8° to 278.6° F). The Sodium salt is a white powder very soluble in Water. Under the name of **Thermiol** a 25 p.c. solution of this salt has been introduced into medicine. It is used as an inhalation in the form of 1 to 3 p.c. solution for tuberculosis and affections of the throat and lungs

Not Official

SODII OLEATIS SOLUTIO, *see* p 824

SODII ET POTASSII TARTRAS

See SODA TARTARATA, p 1088

SODII PHOSPHAS.

SODIUM PHOSPHATE

$\text{Na}_2\text{HPO}_4, 12\text{H}_2\text{O}$, eq 355.64

FR., PHOSPHATE MONO ACIDE DE SODIUM, GER., Natriumphosphat, ITAL., FOSFATO BISODICO, SPAN., FOSFATO SODICO

Colourless, translucent, efflorescent, rhombic crystals, possessing a cooling taste and alkaline reaction

There are three Sodium Phosphates, the Ortho-, Meta-, and Para-phosphates. The official salt is the Di sodium Hydrogen Ortho phosphate

It should be kept in well-closed vessels and in a cool atmosphere, as it has a tendency to effloresce on exposure to air. It may be prepared by the interaction

of Acid Calcium Phosphate with Sodium Carbonate Acid Calcium Phosphate is produced on mixing Bone-ash and Sulphuric Acid

The exsiccated salt, **Sodii Phosphas Exsiccatus**, forms an odourless white powder, which is convenient for mixing with other powders 1 of the dried salt equals about 2 of the crystalline

It is liable to be contaminated with Arsenic of course only Arsenic free samples should be used in medicine

Solubility—1 in 6 of Water, dissolves in its own Water of crystallisation below 212°F , insoluble in Alcohol (90 p c)

Medicinal Properties.—A mild, saline purgative, from its pure saline taste it is called Tasteless Aperient Salt, and is often given to children Diuretic, antacid and antilithic in small doses As it renders the urine alkaline, it is sometimes useful in gout

In a case of diabetes mellitus, 20 grains twice daily by the mouth, a solution being subsequently used hypodermically —*B M J* '03, i 1205

By hypodermic injection in various nervous diseases —*B M J E* '93, ii 108
Incompatible with alkaloids —*T G* '94, 334

Dose—30 to 120 grains = 2 to 8 grammes, for repeated administration, for a single administration, $\frac{1}{4}$ to $\frac{1}{2}$ an oz = 7.1 to 14.2 grammes

Official Preparation—**Sodii Phosphas Effervesceus** Used in the preparation of **Ferri Phosphas**

Not Official—**Liquor Sodii Phosphatis Compositus**

Foreign Pharmacopœias—Official in all except Dan and Norw

Sodii Phosphas Exsiccatus is official in Swiss and U S

Tests—Sodium Phosphate when exposed to dry air loses, according to the *U S P*, 5 molecules of Water of crystallisation, equivalent to 25.1 p c, the *B P* states that when heated to a dull red heat it loses its Water of crystallisation, equivalent to 62.84 of its weight, the *U S P* states that at 100°C (212°F) it loses all its Water of crystallisation, equivalent to 60.3 p c and that at a red heat it is converted into Sodium Pyrophosphate The crystallised salt liquefies at about 40°C (104°F) It answers the tests distinctive of Sodium given under that heading It dissolves readily in Water, yielding a clear solution which is slightly alkaline in reaction towards red Litmus paper The aqueous solution yields on the addition of Silver Ammonio-nitrate Solution a light yellow precipitate readily soluble in Ammonia Solution, and in cold diluted Nitric Acid Magnesium Ammonio-sulphate Solution yields with the aqueous solution a white crystalline precipitate soluble in diluted mineral acid The aqueous solution containing some free Nitric Acid affords on warming with an excess of Ammonium Molybdate Solution a yellow precipitate soluble in Ammonia and reprecipitated as a white crystalline precipitate on the addition of Magnesium Ammonio-sulphate Solution No requisite percentage of pure crystallised Sodium Phosphate is mentioned in the *B P*, nor is a method of determination included The percentage of pure crystallised Sodium Phosphate present may be determined by direct titration with Normal Volumetric Sulphuric Acid Solution, employing Methyl Orange Solution as an indicator of neutrality as described under **Sodii Arsenas**.

On account of the high molecular weight of the salt, 3 grammes is suggested as a suitable quantity to be used for the determination. The *USP* requires that the uneffloresced salt should contain not less than 99 per cent of pure Di-sodium Ortho phosphate, but gives no method of determination. The *PG* does not state either a requisite percentage or a method of determination.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Zinc, Ammonium, Calcium, Potassium, Carbonates, Chlorides and Sulphates. The discovery of Arsenic in the commercial salt caused a sensation in 1900, and numerous methods were suggested with a view to arriving at a satisfactory test for Arsenic. Its presence may be detected by the modified Gutzet's test mentioned below, which is that employed by the *USP*. The *PG* employs the Bettendorf's test given under the heading of Stannous Chloride. Standards have been suggested (*CD* '08, 1796) of 5 parts per million for Lead, and 5 parts per million for Arsenic. Copper, Lead, Iron and Zinc, if present, may be detected by Hydrogen Sulphide, either in a solution rendered acid by Diluted Hydrochloric Acid or in a solution rendered ammoniacal by Ammonia Solution, as described under the heading of Hydrogen Sulphide in small type below. An aqueous solution of the salt should not afford an ammoniacal odour when boiled with Potassium Hydroxide Solution, nor should the issuing vapour possess an alkaline reaction towards a piece of moistened red Litmus paper, indicating the absence of Ammonium salts. The aqueous solution should not afford a distinct opalescence with Ammonium Oxalate Solution, indicating the absence of Calcium. It should not impart a decided violet coloration to a non luminous flame when viewed through a piece of blue glass, indicating the absence of Potassium. It should yield no effervescence on the addition of Diluted Sulphuric Acid, indicating the absence of Carbonate. The aqueous solution when acidified with Nitric Acid should yield not more than a slight reaction for Chlorides and Sulphates when examined by the tests described below under the headings of Silver Nitrate and Barium Nitrate.

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by T.S. of Hydrogen Sulphide, *PG*, slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *USP*.

Barium Nitrate—The aqueous solution (1-20) acidulated with Nitric Acid should not be rendered turbid more than opalescent with Barium Nitrate T.S. within 3 minutes, *PG*.

Silver Nitrate—An acidulated solution as above should not be rendered more than opalescent within 3 minutes by T.S. of Silver Nitrate, *PG*.

Gutzet's Test—5 c.c. of an aqueous solution of the salt (1:10) should not respond to the modified Gutzet's test for Arsenic, *USP*.

Stannous Chloride—A mixture of 1 gramme of previously dehydrated and powdered salt and 3 c.c. of Stannous Chloride T.S. should not assume a dark colour in the course of an hour, *PG*.

Preparation

SODII PHOSPHAS EFFERVESCENS **EFFERVESCENT SODIUM PHOSPHATE**

Sodium Phosphate, in crystals, 50, Sodium Bicarbonate, in

powder, 50, Tartaric Acid, in powder, 27, Citric Acid, in powder, 18, made into granules, the total weight of which is about 100

Dose—60 to 120 grains = 4 to 8 grammes, for repeated administration, for a single administration, $\frac{1}{4}$ to $\frac{1}{2}$ an oz = 7 1 to 14 2 grammes

Official in U S

Not Official

LIQUOR SODII PHOSPHATIS COMPOSITUS—Sodium Phosphate, 100, Sodium Nitrate, 4, Citric Acid, 18, Distilled Water, *q s* to make 100 Triturate the Sodium Phosphate and Sodium Nitrate with the Citric Acid until completely liquefied, then add sufficient Water to produce 100 Filter the liquid and keep it in well-stoppered bottles in a moderately warm place—*U S P*

This has been incorporated in the *B P C*

Not Official

SODII PHOSPHAS ACIDUS

Large, translucent, rhombic crystals, readily soluble in Water Given in cases of alkaline urine

Results showing the power of the drug to increase the acidity of the urine—*B M J* '03, 1 1256, *L* '03, 1 662

Dose—30 to 60 grains = 2 to 4 grammes every 3 hours, but it is better to give smaller quantities oftener to ensure continuous elimination

Sodii Pyrophosphas in Swiss and U S

SODII SALICYLAS.

SODIUM SALICYLATE

$\text{NaC}_7\text{H}_5\text{O}_2$, eq 158 89

FR, SALICYLATE NEUTRE DE SODIUM, GER, NATRIUMSALICYLAT, ITAL, SALICILATO DI SODIO, SPAN, SALICILATO SODICO

There are two Sodium Salicylates, the one prepared with the 'Natural' Acid, the other with the 'Artificial' Acid

The former is in yellowish or pinkish-white pearly scales, or as a pinkish-white amorphous powder, possesses a sweetish saline taste, and frequently a faint odour of Methyl Salicylate, the latter is in white lustrous pearly scales, or a white amorphous powder, with a sweetish saline taste They may be obtained by the interaction of the respective Salicylic Acids and Sodium Carbonate or Sodium Hydroxide

Both varieties should be kept in well-closed bottles of a dark amber tint

The *B P* formula for the salt shows $\frac{1}{2}$ a molecule of Water of crystallisation The *U S P* formula represents the salt as anhydrous, which is correct 100 parts of Sodium Salicylate contain 86 parts of Acid Salicylic

Solubility.—1 in 1 of Water, 1 in 5 of Alcohol (90 p c), 1 in 30 of Absolute Alcohol

Medicinal Properties.—Given as a specific in acute rheumatism in which it lowers the temperature, lessens the pain and swelling, and also the liability to complications such as pericarditis

Occasionally used as an antipyretic in pneumonia, typhoid and all pyrexial affections. A soluble form of Salicylic Acid, and less irritating. Useful in influenza, diabetes, chronic rheumatism, sciatica and in acute tonsillitis, which is so often rheumatic in origin. One of the best antiseptics for fermentative dyspepsia. It increases the acidity of the urine. Brunton says that in obstinate constipation due to gout its administration will tend to keep the bowels regular without any purgative whatever.

Combined with Potassium Bromide, in headache, *P*, *li* 101, *T G* '94, 935, in pleuritis, *T G* '94, 101, reason for advantage of natural over artificial Salicylate, *P*, *li* 1 447, of great value in pneumonia and in many forms of erythema, especially *e. nodosum*, *L* '80, 1 627 '95, 1 1422, *B M J* '86, 1 737, *T G* '85, 446. In exophthalmic goitre — *B M J E* '95, 1 91.

As a means of diagnosis between rheumatism and gout, if the patient improved under Salicylate treatment the disease was rheumatic, if not it was gout — *L* '99, 1 441.

Larger doses of the salt prepared from the 'natural' acid could be given with less ill effects — *L* '00, 1 1016.

In pneumonia 8 to 10 grains every 2 hours — *Pr* *lxiv* 380.

Temporary blindness resulting from 140 to 150 grains taken over a period of 60 hours — *B M J* '01, 1 81.

10 to 20 grains combined with 10 grains Quinine Sulphate every 4 hours in malarial fever — *L* '03, 1 95, 200, 681.

In the treatment of chorea 10 grain doses with 20 grains Sodium Bicarbonate for a child of 6 to 10 years, increasing the quantity to 15 and 30 grains respectively, after 2 or 3 days and if necessary to 20 and 40 grains respectively after a further 2 or 3 days. A careful watch is kept for any symptoms of Salicylate poisoning — *B M J* '03, 1 451.

Has been used as an endo-articular and as an intravenous injection in sterile Water, under strict aseptic precautions, in cases of acute articular rheumatism. *M P* '04, 1 472, *B M J E* '04, 1 60.

Intravenous injection of 2 c.c. of Mendel's mixture, the latter containing 17.5 p.c. of Sodium Salicylate and 2.5 p.c. Caffeine, in the treatment of rheumatic affections. When injected into the veins one sees the specific action of Salicylates in rheumatism at its best — *B M J E* '05, 1 43.

Very beneficial in posterior urethritis — *F T* '07, 84. No need to discontinue it during pericarditis — *B M J* '07, 1 814.

Dose — 10 to 30 grains = 0.65 to 2 grammes.

Prescribing Notes — *Best given in solution well diluted, to avoid dyspepsia, but may also be prescribed in cachets or powders. When dissolved in Water and mixed with Ammonia, the solution soon becomes yellow or brown on exposure to the air, which happens in mixtures containing the salt and Aromatic Spirit of Ammonia when the bottle is half full. It is sometimes prescribed with Citric Acid, which precipitates the Salicylic Acid. It is better to give it with Sodium or Potassium Citrate. When prescribed with a salt of Quinine, Quinine Salicylate is formed, which is only slightly soluble, and is therefore thrown out.*

Official Preparation — Used in the preparation of Bismuthi Salicylas.

Foreign Pharmacopœias — Official in all the Foreign Pharmacopœias except Port. Dutch has also Salicylas Natricus cum Coffeino.

Tests — Sodium Salicylate when heated emits white inflammable vapours, possessing an odour of Phenol, leaving a carbonaceous residue, which, when dissolved in Water, produces a solution having a strong alkaline reaction towards red Litmus paper, and which effervesces on the addition of a diluted mineral Acid. It dissolves readily in Water, forming a clear solution which is neutral to Litmus paper or only faintly acid towards blue Litmus paper. The *B P*

states that a concentrated aqueous solution affords with Ferric Chloride T S a reddish-brown coloration, and a diluted aqueous solution a violet coloration with the same reagent. The *USP* states that Ferric Chloride T S added to an excess of a concentrated aqueous solution of the salt produces a violet precipitate, but when added to a diluted solution (1 in 100) it produces a deep violet-blue colour, the *PG* states that a 1 in 1000 aqueous solution affords a bluish-violet coloration on the addition of Ferric Chloride T S. A 1 in 20 aqueous solution affords a green coloration with Copper Sulphate T S. A 10 p c aqueous solution yields on the addition of Diluted Sulphuric Acid a white crystalline precipitate readily soluble in Ether. If the precipitate produced on acidification be separated by filtration, washed and carefully dried, it should possess the m p and answer the tests distinctive of Salicylic Acid given under Acidum Salicylicum. A small quantity of the salt, when warmed with a little concentrated Sulphuric Acid and a few drops of Methyl Alcohol, evolves a distinctive odour of Methyl Salicylate. Neither the *BP* nor the *PG* states what percentage of pure Sodium Salicylate should be present in the salt, nor does either give a method of determination. It may be determined by titration of the solution of the residue left on ignition, with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as an indicator of neutrality, 1 c c of the Normal Volumetric Acid Solution being equivalent to 0.15889 gramme of pure Sodium Salicylate. The *USP* requires it to contain 99.5 p c of pure Sodium Salicylate as volumetrically determined by titrating the solution obtained by exhausting the residue left on thoroughly igniting the salt at a red heat, with boiling Water, until the washings cease to react with Methyl Orange Solution as described in small type below under the heading Volumetric Determination.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Zinc, Chlorides, Sulphates and Sulphites, organic impurities and Carbonates, unconverted Phenol, and isomers or homologues of Salicylic Acid. Copper, Lead, Iron and Zinc, if present, are indicated by the time-limit test described under the heading of Hydrogen Sulphide. When Diluted Nitric Acid is added in slight excess to an aqueous solution of the salt and the precipitated Salicylic Acid is removed by filtration, the filtrate should yield only a slight turbidity on the addition of Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than traces of Chlorides and Sulphates. The *BP* states that if the aqueous solution be acidulated with Nitric Acid and the precipitate be dissolved with a little Alcohol (90 p c) the mixture affords not more than the slightest reactions with the tests for Sulphates or for Chlorides. The word 'acidulated' should be 'supersaturated,' as sufficient Nitric Acid over and above that necessary to completely decompose the Sodium Salicylate must be added in order to prevent the precipitation of Silver Salicylate. If a 5 p c aqueous solution of the salt be mixed with a few drops of Iodine T S and a few drops of Hydrochloric Acid, the filtrate should yield no precipitate upon the addition of

Barium Chloride Solution, indicating the absence of Sulphites. The salt should dissolve without coloration and without effervescence in cold Sulphuric Acid, indicating the absence of organic impurities and of Carbonates. The concentrated aqueous solution, when shaken with an equal volume of Ether, and the ethereal solution allowed to evaporate spontaneously, the residue should be free from any odour of Phenol. The *BP* states that 50 to 100 grammes of the salt, kept in a closed vessel for several days, should not evolve the faintest odour of Phenol. Isomers and homologues of Salicylic Acid can be detected, if present, by their influence on the m.p. of the acid separated from the salt on acidification. The *BP* includes a test for distinguishing Salicylates from Carbates and Sulphocarbates. A solution containing not less than 1 p.c. is stated to afford a yellowish-brown precipitate with Uranium Nitrate Solution. Carbates and Sulphocarbates presumably afford no precipitate with Uranium Nitrate Solution, it is difficult, therefore, to gauge the value of this test, as it will certainly not detect the presence of Carbates and Sulphocarbates in the Salicylate. The test also appears under Acidum Salicylicum, and is commented upon in large type under the heading of Tests.

Sulphuric Acid—The salt is soluble without effervescence or coloration in cold Sulphuric Acid, *BP* and *PG*, the *PG* uses 0.1 gramme of salt and 1 c.c. of acid.

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by T.S. of Hydrogen Sulphide, *PG* acidulated with Hydrochloric Acid and filtered, the filtrate should not respond to the time limit test for heavy metals, *USP*.

Silver Nitrate—2 volumes of an aqueous solution (1-20) mixed with 3 volumes of Alcohol (90 p.c.) and acidified with Nitric Acid should not be affected by T.S. of Silver Nitrate, *PG*. Also given in *BP* without quantities.

Barium Nitrate—An aqueous solution (1-20) should not be affected by T.S. of Barium Nitrate, *PG*. The *BP* directs the addition of Nitric Acid and Alcohol, as in the Silver Nitrate test, and uses Barium Chloride Solution.

Iodine and Barium Chloride—If to an aqueous solution of the salt (1-20) 3 drops of Iodine T.S. and a slight excess of Hydrochloric Acid be added, the filtrate from this mixture should not yield a precipitate upon the addition of T.S. of Barium Chloride, *USP*.

Volumetric Determination—If 1 gramme of the dry salt be thoroughly ignited at a red heat, and the residue extracted with boiling Distilled Water until the washings cease to react with Methyl Orange T.S., the mixed filtrate and washings should require for complete neutralisation not less than 12.5 (12.52) c.c. of Semi-normal Volumetric Sulphuric Acid Solution, Methyl Orange T.S. being used as indicator, *USP*.

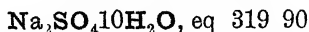
Not Official

SODII DITHIO-SALICYLAS (Dithion)—A yellowish white, amorphous, somewhat hygroscopic powder, antiseptic and antipyretic. Used in the form of powder, solution or ointment. Has been found useful in the treatment of rheumatism.

Dose—1 to 3 grains = 0.06 to 0.2 gramme

Liquor Natri Silicici (Soluble Glass) is official in Austri and Ger.

SODIUM SULPHATE.



FP , SULFATE DE SODIUM OFFICINAL, GER , NATRIUMSULFAT ,
ITAL , SOLFATO DI SODIO , SPAN , SULFATO SODICO

It should be kept in well-closed vessels and in a cool atmosphere, as it is readily effloresced on exposure to air, losing its Water of crystallisation.

Sodii Sulphas Exsiccatus, is an odourless white powder, 1 of which equals $2\frac{1}{2}$ of the crystalline salt. Much more convenient than the latter in mixing with other powders.

Solubility—1 in 3 of Water, and measures 3½, 10 in 3 of Water at 92° F, 10 in 4½ of Water at 212° F, insoluble in Alcohol (90 p c)

Medicinal Properties --Hydragogue purgative and cholagogue, useful in cases of gall-stones and of liver disease, in small repeated doses it is especially well adapted for cases of constipation associated with gout and hepatic dyspepsia

Given in 1-drm doses in either Fennel or Cinnamon Water 4, 5 or 6 times a day in the treatment of dysentery — *I M G* '05, n 280 In acute cases no drug is known which acts so rapidly, painlessly, or so effectually

Dose—30 to 120 grains = 2 to 8 grammes, for repeated administration, for a single administration, $\frac{1}{4}$ to $\frac{1}{2}$ an ounce = 7 1 to 14 2 grammes

Official Preparation —Sodii Sulphas Effervescens

Not Official—Pulvis Sodii Sulphatis et Zingiberis, Pulvis Salis Carolini
Factitii Effervescens, Sal Carolinum Factitium

Foreign Pharmacopœias—Official in Hung (*Natrium Sulfuricum Crystallissimum*), also *Siccum*, Dan, Dutch, Norw and Swed (*Sulphæs Nitricus*), Dan and Swed, also *Siccatus*, Dutch, also *Exsiccatus*, Fr. (*Sulfate de Sodium Officiel*), Aust, Belg, Jap and Swiss (*Natrium Sulphuricum*), also *Siccum*, Ital (*Solfato di Sodio*), Mex (*Sulfato de Sodio*), Port (*Sulphato de Soda*), Russ (*Natrium Sulfuricum*), Depuratum, and *Siccum*, Span (*Sulfato Sodico*), U.S.

Tests—Sodium Sulphate melts when heated. The *U.S.P.* says the salt fuses at 33° C (91 4° F). When dried at 100° C (212° F) it loses the whole of the Water of crystallisation, equivalent to 55.9 p.c., the *B.P.* states when exposed to heat in a porcelain crucible, but gives no indication as to the temperature. It answers the tests distinctive of Sodium given under that heading. It dissolves readily in Water, forming a clear solution which is neutral in reaction towards Litmus paper, and which, on the addition of Barium Chloride Solution, yields a white precipitate insoluble in Hydrochloric Acid. It is officially required to contain 100.0 p.c. of pure crystallised Sodium Sulphate, as gravimetrically determined by BaCl_2 . Aqueous solution of 1 gramme of the salt, acidulated with Hydrochloric Acid, with Barium Chloride Solution, the white precipitate produced, when well washed and dried, should weigh 0.725 of a gramme. The *U.S.P.*

requires the salt to contain in an uneffloresced condition not less than 99 p.c. of pure crystallised Sodium Sulphate, but no method of determination is given. The *P G* does not state either a percentage or a method of determination.

The more generally occurring impurities are Arsenic, Copper, Lead and Iron, Ammonium, Calcium, Magnesium and Potassium, Carbonates and Chlorides. The *B P* includes also a test for Aluminium, an impurity of comparatively trivial importance, but omits a test for Arsenic, a much more likely one to be found and one possessing much greater importance. The presence of Arsenic much in excess of 1 in 100,000 is indicated by the sample responding to the modified Gutzeit's test described below. The *P G* employs the Bettendorf test for Arsenic with Stannous Chloride Solution. Standards have been suggested (*C D* '08, 1796) of 5 parts per million for Lead, and 2 parts per million for Arsenic. Copper, Iron and Lead, if present, may be indicated by the coloration produced by Hydrogen Sulphide in either a solution slightly acidified with Hydrochloric Acid or a solution made slightly alkaline with Ammonia. The *P G* includes a separate test for Iron described below under the heading of Potassium Ferrocyanide. The aqueous solution of the salt should not afford an ammoniacal odour when boiled with Liquor Potassæ, nor should the issuing gas possess an alkaline reaction towards moistened red Litmus paper, indicating the absence of Ammonium salts. It should yield no turbidity with Ammonium Oxalate Solution, indicating the absence of Calcium, and when to the mixture is added Ammonium Chloride, allowed to stand some time and filtered, the filtrate should not yield a turbidity on the addition of Sodium Phosphate Solution, indicating the absence of Magnesium. When viewed through a piece of blue glass no violet coloration should be imparted to a non-luminous flame when a crystal of the salt moistened with Hydrochloric Acid is introduced, indicating the absence of Potassium. A strong aqueous solution of the salt should yield no effervescence on the addition of Hydrochloric Acid, indicating the absence of Carbonates. When acidified with diluted Nitric Acid it should yield no decided turbidity on the addition of Silver Nitrate Solution, indicating the absence of more than traces of Chlorides. Excess of moisture may be detected by a loss on drying at 100° C (212° F) as described above.

Hydrogen Sulphide—An aqueous solution (1-20) should not be affected by T.S. of Hydrogen Sulphide, *P G*, slightly acidulated with Hydrochloric Acid should not respond to the time limit test for heavy metals, *U S P*.

Sodium Phosphate—An aqueous solution (1-20), after the addition of Ammonia T.S. should not be affected by T.S. of Sodium Phosphate, *P G*.

Silver Nitrate—An aqueous solution (1-20) should not undergo any change within 5 minutes on the addition of T.S. of Silver Nitrate, *P G*.

Potassium Ferrocyanide—20 c.c. of an aqueous solution (1-20) should not be affected by 0.5 c.c. of T.S. of Potassium Ferrocyanide, *P G*.

Gutzeit's Test—5 c.c. of the aqueous solution (1-10) should not respond to the modified Gutzeit's test for Arsenic, *U S P*.

Stannous Chloride—A mixture of 1 gramme of previously dried and

powdered Sodium Sulphate, and 3 c c of Stannous Chloride T S should not assume a dark colour in the course of an hour, *P G*

Preparation

SODII SULPHAS EFFERVESCENS EFFERVESCENT SODIUM SULPHATE

Sodium Sulphate, in crystals, 50, Sodium Bicarbonate, in powder, 50, Tartaric Acid, in powder, 27, Citric Acid, in powder, 18, made into granules, the total weight of which is about 100 (1 in 2)

Dose—60 to 120 grains = 4 to 8 grammes, for repeated administration, for a single administration, $\frac{1}{4}$ to $\frac{1}{2}$ an ounce = 7 1 to 14 2 grammes

Not Official

PULVIS SODII SULPHATIS ET ZINGIBERIS—Sodium Sulphate, in powder, 60 grains, Ginger, in powder, 5 grains, mix

To be taken in a small tumbler of warm Water, in the morning

PULVIS SALIS CAROLINI FACTITII EFFERVESCENS (Effervescent Powder of Carlsbad Salt)—Dried Sodium Sulphate, 11 oz, Powdered Potassium Sulphate, $\frac{1}{2}$ oz, Sodium Chloride, 4 $\frac{1}{2}$ oz, Sodium Bicarbonate, 51 oz, Tartaric Acid, 40 oz, Gluside, 28 grains Dry separately, reduce to fine powder and mix —*B P C Formulary* 1901

Dose—60 to 120 grains = 4 to 8 grammes

Exsiccated Sodium Sulphate, 9, Sodium Potassium Tartarate, 38, Sodium Chloride 3, Sodium Bicarbonate, 33, Gluside 0 05, Tartaric Acid, *q s* to produce 100 —*B P C*

SAL CAROLINUM FACTITIUM—Dry Sodium Sulphate, 22, Potassium Sulphate, 1, Sodium Chloride, 9, Sodium Bicarbonate, 18 —*Ger*

This has been incorporated in the *B P C*

Dry Sodium Sulphate 47, Potassium Sulphate 2, Sodium Chloride 15, Sodium Bicarbonate 36 —*Jap*

SODII SULPHIS.

SODIUM SULPHITE

$\text{Na}_2\text{SO}_3 \cdot 7\text{H}_2\text{O}$, eq 250 38

Colourless, transparent, efflorescent, monoclinic prisms, having a cooling saline and sulphurous taste It can be prepared by interaction of Sulphurous Acid and Sodium Carbonate

It should be preserved in well-closed bottles and kept in a cool place, as both the crystals and aqueous solutions are liable to oxidation on exposure to air

Solubility—3 in 4 of Water, insoluble in Alcohol (90 p c), 1 in 25 of Glycerin

Medicinal Properties.—Antiseptic, given with success in fermentative vomiting and dilated stomach due to *sarcina ventriculi* Externally as a lotion in parasitic cutaneous affections.

Dose.—5 to 20 grains = 0 32 to 1 3 gramme,

Incompatibles—II a.c. Acids,

Not Official—Liquor Sodii Sulphatis Benzoicus, Sodium Thiosulphate and Lotio Sodii Hyposulphitis

Foreign Pharmacopœias—Official in Mex., Port. and U.S.

Tests—Sodium Sulphate when gently heated gradually loses its Water of crystallisation, and at a temperature of a little above 100°C (212°F) it loses its Water of crystallisation, equivalent to 50 p.c. It answers the tests distinctive of Sodium given under that heading. It dissolves readily in Water, yielding a solution which is usually faintly alkaline towards red Litmus paper. The *USP* states that it is neutral or feebly alkaline to Litmus paper. It evolves on the addition of Hydrochloric Acid a colourless gas possessing the characteristic pungent odour of burning Sulphur. On the addition of Zinc and Hydrochloric Acid it evolves the characteristic odour of Hydrogen Sulphide, and if a piece of filter paper moistened with Lead Acetate Solution be suspended in the neck of the tube it acquires a black colour. Iodine Solution added to an acidified solution is instantly decolorised. It is officially required to contain not less than 97.3 nor more than 102.3 p.c. of pure crystallised Sodium Sulphate as volumetrically determined by the method given below under the heading of Volumetric Determination. The *USP* requires that it should contain in the uneffloresced and anhydrous condition not less than 94 p.c. of pure Sodium Sulphate as volumetrically determined by the method also given below in small type under the heading of Volumetric Determination. The salt is not official in the *PG*.

The more generally occurring impurities are Arsenic, Copper, Lead and Iron, Chlorides, Sulphates and Thiosulphates. The *BP* includes only a test for the latter, the *USP* tests for the heavy metals and Thiosulphates. Arsenic, Copper, Lead and Iron, if present, may be detected by the Hydrogen Sulphide test described below. The strongly acidified aqueous solution of the salt should yield no decided turbidity on the addition of Silver Nitrate T.S., indicating the absence of more than traces of Chlorides. It should not yield a pronounced turbidity on the addition of Barium Chloride Solution to an aqueous solution strongly acidified with Hydrochloric Acid, indicating the absence of more than traces of Sulphates. The aqueous solution when treated with Hydrochloric Acid should not become cloudy, indicating the absence of Thiosulphate. The *USP* employs diluted Nitric Acid as a test for the absence of Thiosulphate, when heated sufficiently to expel the gases, no turbidity should appear.

Time-limit Test—A solution of 1 gramme in 20 c.c. of Diluted Hydrochloric Acid after heating sufficiently to expel the Sulphur Dioxide and restoring the solution to its original volume, should not respond to the time limit test for heavy metals, *USP*.

Volumetric Determination—A solution of 1 gramme of the salt in 50 c.c. of Water should decolorise not less than 77.7 nor more than 81.7 c.c. of Volumetric Solution of Iodine, *BP*. If to 50 c.c. of Tenth normal Iodine Volumetric Solution measured from a burette into a glass stoppered vial (of about 100 c.c. capacity), 0.5 gramme of the finely powdered crystals of Sodium Sulphate be added, after solution has taken place, not more than 12.45 c.c. of Tenth normal Volumetric Sodium Thiosulphate Solution should be required to discharge the colour of the solution, *USP*.

Not Official

LIQUOR SODII SULPHITIS BENZOICUS — Sodium Sulphite, 30, Benzoic Acid, 14, Water, 500 An Antiseptic solution, recommended by Heckel

SODIUM THIOSULPHATE (Sodium Hyposulphite $\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$, eq 246.44) — Colourless transparent monoclinic prisms, possessing a cooling and somewhat bitter, sulphurous taste Soluble 5 in 3 of Water, insoluble in Alcohol (90 p c) It is seldom used internally as a medicinal agent, but on account of its poisonous influence on the sacina ventriculi which attends yeast vomiting it has been employed in that complaint Externally in the form of a 12½ p c solution it has been used in parasitic diseases It is used for removal of stains produced by Silver salts and in volumetric analysis

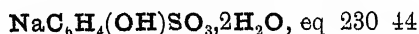
Dose — 5 to 10 grains = 0.32 to 0.65 gramme

Foreign Pharmacopœias — Official in Fr, Ger, Swiss and U S

Lotio Sodii Hyposulphitis — Sodium Hyposulphite, 1 dram, Water, to 1 fl oz — *St John's*

SODII SULPHOCARBOLAS.

SODIUM SULPHOCARBOLATE



Colourless, translucent, slightly efflorescent rhombic crystals, possessing at first a saline and subsequently a slightly bitter taste It may be prepared from Para-phenol-sulphonic Acid obtained by Phenol in excess of Sulphuric Acid, by converting it salt

It should be kept in well-stoppered glass bottles and in a cool atmosphere, as it has a tendency to effloresce in dry air

The Sulphocarbolates used in medicine are defined as the salts of Para-phenol-sulphonic Acid The action of Sulphuric Acid upon Carbohic Acid results in a mixture of Para- and Ortho phenol-sulphonic Acids, the proportion of the latter being less the higher the temperature, and the longer continued the contact To eliminate the Ortho salt further purification is necessary

Solubility — 1 in 6 of Water, 1 in 150 of Alcohol (90 p c), 1 in 5½ of Glycerin

Medicinal Properties — Antiseptic, given in cases of flatulence, fermentative dyspepsia, and other conditions in which Carbohic Acid is used

Dose. — 3 to 15 grains = 0.2 to 1 gramme

Incompatibles — Ferric salts

Foreign Pharmacopœias — Official in Jap and U S (Sodii Phenol sulphonas)

Tests. — Sodium Sulphocarbolate when heated at a temperature slightly above 100° C (212° F) loses its Water of crystallisation, equivalent to 15.5 p c When more strongly heated it evolves inflammable vapours possessing a characteristic odour of Phenol, and when ignited leaves a residue of Sodium Sulphate, equivalent to 30.6 p c of its original weight This residue answers the tests distinctive of Sodium given under that heading, and when dissolved in Water yields with Barium Chloride Solution a white

precipitate insoluble in Hydrochloric Acid. The salt dissolves readily in Water, yielding a solution which is neutral in reaction towards Litmus paper. A diluted aqueous solution of the salt yields on the addition of Ferric Chloride T.S. a violet coloration. The *B.P.* does not state what percentage of the pure salt it should contain, nor does it include a method of determination. The *U.S.P.* requires that it should contain not less than 99 p.c. of pure crystallised Sodium Para phenol-sulphonate, and, although it does not indicate a direct determination to be made, states that the residue of Sodium Sulphate left on ignition should amount to 30.6 p.c.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Sulphates. The presence of Arsenic may be ascertained by the modified Gutzeit's test or by the test with Hydrogen Sulphide described below. The latter test when applied either in solution slightly acidified with diluted Hydrochloric Acid or in a solution rendered alkaline with Ammonia, serving to detect also Copper, Lead and Iron. The aqueous solution of the salt should not at once be rendered turbid by Barium Chloride Solution, indicating the absence of Sulphates. The *U.S.P.* states that a diluted solution of the salt (1 in 100) remains clear on the addition of Barium Chloride T.S. The *B.P.* states that it may be distinguished from Salicylate by not yielding a yellowish-brown precipitate with Uranium Nitrate Solution. The criticisms on this latter test will be found under the headings Acidum Salicylicum, Sodii Salicylas.

Time-limit Test—The aqueous solution of the salt (1-20), slightly acidulated with Hydrochloric Acid, should not respond to the time limit test for heavy metals, *U.S.P.*

Not Official

SODII SULPHOVINAS

SODIUM SULPHATHYLATE

Translucent, hexagonal crystals, or as a white, granular powder, very hygroscopic, and should be kept in well stoppered bottles. It is soluble in Water, in dilute Alcohol and in Glycerin. Used as a mild aperient.

Dose— $\frac{1}{2}$ to 1 oz. = 14.2 to 28.4 grammes.

Not Official

SODII TAUROCHOLAS

A yellow or yellowish brown, amorphous, granular, hygroscopic powder, or a brown or blackish brown, sticky, resinous mass. Soluble 2 in 1 of Water, partially soluble in Alcohol (90 p.c.). It is best prepared from pig's bile.

It should be kept in well closed bottles and in a cool place.

Given in gouty obesity and dyspepsia, 4 grains immediately after each meal. The pills should be coated with Keratin.

Dose—2 to 6 grains = 0.13 to 0.4 gramme made into pill with Alcohol (60 p.c.).

Tests—Sodium Taurocholate on ignition yields a carbonaceous residue, possessing, when dissolved in Water, a strong alkaline reaction towards red Litmus paper, and effervesces on the addition of Hydrochloric Acid. It answers the tests distinctive of Sodium given under that heading. The salt dissolves

readily in Water, yielding a solution which affords a precipitate on the addition of Ammonia Solution and basic Lead Acetate. Taurocholic Acid may be determined by the amount of Sulphur which it contains. A weighed quantity of the salt is moistened with fuming Nitric Acid and evaporated to dryness on a water-bath; the residue is dissolved in Water, the solution filtered and precipitated by the addition of Barium Chloride Solution when the precipitate is filtered off, washed, dried, ignited and weighed, 1 part of Barium Sulphate corresponding to 2.16 parts of Taurocholic Acid.

Acidum Taurocholicum (Taurocholic Acid) forms deliquescent silky needles, readily soluble in Water and in Alcohol (90 p.c.)

Sodii Glycocholas occurs together with the above salt in ox bile. It may be obtained in the form of stellate needles. It is given in gouty obesity and dyspepsia, and has been found to possess considerable cholagogue action.

Dose — 2 to 10 grains = 0.13 to 0.65 gramme

Not Official

SODII VANADAS

White or yellowish white, odourless, granular powder, soluble 2 in 1 of Water, insoluble in Alcohol (90 p.c.)

Stimulates the gastric mucosa, increases the appetite and improves the general condition — *B.M.J.* '01, 11 88, *C.D.* '02, 1 638

Dose — $\frac{1}{8}$ to $\frac{1}{2}$ grain = 0.001 to 0.0027 gramme

Iron Meta-vanadate, a dark, greyish-brown powder, insoluble in Water and in Alcohol (90 p.c.), and **Lithium Meta-vanadate**, a yellowish-white crystalline powder, soluble in Water, have also been prepared.

SOLUBILITY

The importance of the subject of solubility to medical men was recognised as far back as 1864, by the late Peter Squire, and consequently this has been a feature of Squire's *Companion* to the *B.P.* since the first edition. The several paragraphs under this title are probably of more use to the prescriber and the pharmacist than others in a book of this kind. The prescriber is constantly wishing to know how much of a given substance will dissolve in some liquid which he desires to use, and to what extent. It is obvious that an error stating the substance to be more soluble than it really is causes more trouble and error in the opposite direction, but the figures should at least be correct. Prior to 1885 very few substances were given in the *B.P.*, but subsequently these were very much enlarged.

Figures for the solubility of the various substances have been given in the *Companion* since its first issue in 1864, and these have been revised and supplemented from time to time in subsequent editions, from experiments made for that purpose. In most instances the figures have been ascertained by adding the solid substance in fine powder to a liquid, and shaking it at intervals during 3 days at a temperature between 58° and 62° F (14.4° and 16.6° C). They represent the weight of a solid in grammes, and the measure of a fluid in c.c. Some liquids are stated to be miscible in all proportions, this has been ascertained by adding to 5 c.c. of one fluid, small quantities of the other fluid, $\frac{1}{10}$ c.c. at first, and afterwards $\frac{1}{2}$ c.c. until 20 c.c. have been added, shaking the mixture after each addition, the temperature of the mixture being kept inside the limits given above. At the instance of the Pharmacopœia Committee of the General Medical Council a large number of experiments were made in the Research Laboratory of the Pharmaceutical Society, with a view to determining the accuracy or otherwise of the solubilities of chemicals mentioned in the British Pharmacopœia, and the results were fully reported and a comparison made with authoritative statements — *P.J.* '00, 11 190, '01, 1 774, 806, '02, 1 510, 532, 551. These reports have been closely criticised and compared with

Squire's *Companion* figures in a series of papers by the author and C M Caines, communicated to the *Pharmaceutical Journal* and to the *Chemist and Druggist*.

Out of a number of determinations, amounting in the aggregate to 91 figures, 74 are almost identical, 12 are within the range of individual experimental error and variations in commercial samples. In 3 cases Squire's *Companion* figures represent the solubility in Alcohol (88.7 p.c.) of the *BP* '85, instead of the Alcohol (90 p.c.) in the *BP* '98, and in the remaining 2 instances the Research laboratory figures are shown to be incorrect—*CD* '02, 11 944, '05 1 783, *PJ* '03, 1 65, '05, 1 720. The concluding report from the Research laboratory on the solubility of chemical substances mentioned in the British Pharmacopœia (*PJ* '08, 11 881, 945) states that the further experiments confirm the results obtained by Squire and Caines. A short reference also appears under the individual heading of each substance, and will be found in the large type under Ammonii Phosphas and Zinci Sulphocarbolas.

In the Continental Pharmacopœias solubility figures are usually expressed in parts by weight, and this fact is frequently overlooked when such figures are noted by other workers. In the case of liquids lighter or heavier than Water the difference may be considerable. For instance, in a communication from the Research laboratory (*PJ* '03, 11 946), 'A comparison with authoritative statements,' shows an apparent discrepancy between the figures given in the British and German Pharmacopœias for the solubility of Phenacetin in Alcohol (90 p.c.) 1 in 20 and 1 in 16 respectively, whereas the British being given by volume and the German by weight, the figures are in perfect accord. It would have been advisable (*PJ* '05, 1 720) to insert the sp. gr. of the menstruum and the words 'by weight' against the Continental authority. The figures given in the *8th Decennial Revision of the USP* represent those obtained at a temperature of 25° C (77° F). It is stated in the preface that this temperature was adopted for solubilities after much discussion, because it is believed that it will be generally more satisfactory throughout the United States than the former temperature of 15° C (59° F)—the average temperature of laboratories and stores in the United States throughout the year being nearer 25° C (77° F) than 15° C (59° F). The above requires to be carefully noted, more particularly in instances where comparisons are being made between figures appearing in books of recognised standing, and more especially in books of reference where the figures are not the result of actual experiment, but consist of a mere compilation. An instance readily occurs in the case of a recently published text book—it had evidently been held necessary, where figures for the solubility of an individual substance in a certain menstruum were not available in one book, to incorporate figures obtained from another reliable source, and for the accuracy and uniformity of the system the importance of the above caution cannot be overestimated. Thus figures for the solubility of Terpene Hydrate, 1 in 280 of Water, 1 in 14 of Alcohol (90 p.c.), 1 in 46 Alcohol (60 p.c.), 1 in 32 boiling Water, 1 in 2 of boiling Alcohol, 1 in 100 of Ether, and 1 in 200 of Chloroform may be of considerable utility with a definite knowledge that they are the results of determinations made at two different temperatures, but when they are definitely stated all to refer to determinations made at 15.5° C (60° F), whereas in reality they are the result of two different systems, their misleading nature is apparent. The strength of the menstruum used is also a matter of importance. The solubility of some official substances has been shown (*PJ* '05, 1 720) to vary considerably where Alcohol (88.7 p.c.) of *BP* 1885 or Alcohol (90 p.c.) of the *BP* 1898 is used as a menstruum, how much more so then is the solubility likely to vary when Alcohol (94.9 p.c.) of the *USP* is mistaken for Alcohol (90 p.c.) of the *BP*. Again the Ether official in the *BP* refers to Ether, sp. gr. 0.735. Ether (*USP*) refers to an Ether corresponding to the sp. gr. 0.720 of the *BP*, which is another point likely to make a considerable difference, as will be seen by a reference to the note appearing under the solubility of Citric Acid and Tartaric Acid. In addition to substances of official origin, it is frequently necessary to consult figures for the determinations of substances of unofficial origin. A very large number of such determinations have been carried out in the author's laboratory, and the results of these determinations have been incorporated in a series of papers appearing in the pharmaceutical press (*CD* '05, 1 783, *PJ* '05, 1 720, 784).

Not Official

SOMATOSE.

A light, white or greyish powder, stated to be prepared from fresh meat, soluble in Water, and consisting of a mixture of deutero- and hetero Albumoses.

Denaeyer states that it is neither Albumose nor a peptone, but has the characters of an alkali-albumen. This statement is partially confirmed by Allen.

A true meat nutrient, possessing restorative and stimulating powers, being well borne by delicate patients. Has a favourable effect on general metabolism. Produces no irritant effect on the kidneys and it never gives rise to albuminuria, albumosuria or peptonuria.—*L* '99, 11 885

Doses of 12 grains, useful in secondary syphilis, and in the anæmia caused by malaria.—*B M J E* '99, 1 16

Recommended in anæmia, in intestinal disorders, and in dyspepsia.

Liquid Somatose is given in teaspoonful doses to adults.

Iron Somatose—Is a light brown, almost tasteless powder, soluble in aqueous liquids. It contains 2 p.c. of Iron, and has been recommended in chlorosis. **Milk Somatose** has also been introduced.

Not Official

SOZOIODOL

DI-IODOPARAPHENOLSULPHONIC ACID

A white, shining, crystalline powder, containing Iodine about 52 p.c., Carboic Acid 20 p.c., and Sulphur 7 p.c., preferably used in the form of its salts. When required in solution, the Sodium salt is most applicable, dissolving 1 in 14 of Water or Glycerin. The Potassium salt, soluble 1 in 100 of Water, is preferable as a dusting powder, or in ointments. Solution of Zinc salt, 1 to 3 p.c., is suitable for injection.

Medicinal Properties—A substitute for Iodoform.

It is recommended locally in nasal and pharyngeal disorders, and as an antiseptic of great energy in parasitic skin affections.—*B M J* '89, 11 42, *T G* '94, 132, 91, 592. In aural and nasal affections.—*L* '94 1 1636, *B M J E* '94, 1 99

Soziodol cotton and gauze containing 5 and 10 p.c.

HYDRARGYRI SOZOIODOLAS (Mercury Soziodol)—A fine, orange-yellow, amorphous powder, almost insoluble in Water, insoluble in Alcohol (90 p.c.).

It should be kept in well-stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the air and light.

Has been employed in syphilis and in psoriasis, chiefly by hypodermic injection (*see* below).

The injections of this salt are stated to be less painful than those of Mercuric Chloride.—*L* '01, 11 522, '03, 1 785, *B M J* '03, 1 656

Tests—Mercury Soziodolate, although almost insoluble in Water, dissolves in Sodium Chloride Solution. 0.5 of a gramme with 1.5 grammes of Sodium Chloride should dissolve leaving only a faint turbidity. It contains theoretically 32.0 p.c. of metallic Mercury, which may be determined by distillation with Lime. A solution of 0.1 of a gramme in 1 c.c. of Nitric Acid and 9 c.c. of Water should be rendered only faintly turbid on the addition of Silver Nitrate solution, indicating the absence of Chlorides. A solution 0.2 of a gramme of the preparation dissolved in 20 c.c. of Water by the aid of a little Hydrochloric Acid should neither yield a distinct turbidity on the addition of Barium Nitrate Solution, nor on the addition of diluted Sulphuric Acid, and a white precipitate of Sulphates and Barium salts. When ignited with free access of air it should leave no weighable residue.

INJECTIO HYDRARGYRI SOZOIODOLATIS HYPODERMICA —
Mercury Soziodol, 5 grains, Sodium Iodide, 10 grains, Distilled Water,
200 minims Inject 10 to 15 minims = 0.6 to 0.88 c.c. —*Loc'h*

Not Official
S P E R M I N

DR. BROWN SEQUARD'S ORCHIC LULUID

Full details regarding its preparation and uses are published *B M J* '93, 1 1145, 1212, with an editorial article p. 1279, *B M J L* '94, 11 52, 56, *T' G* '93, 110

In the form of an essence, 20 drops taken 3 times a day, in the treatment of abnormalities of frequency and rhythm of the pulse — *L M J E* '02, 1 23, *L* '02, 1 326

SPIRITUS
SPIRIT

All saccharine substances which have undergone the vinous fermentation contain Alcohol, which can be separated by distillation. The various kinds of alcoholic liquids are distinguished by differences in flavour and colour.

When Alcohol is distilled with aromatic substances containing volatile Oil, part of the Oil is carried over by the alcoholic vapour, and condenses along with it.

All the official Spirits, except Brandy, are prepared with Alcohol (90 p.c.)

SPIRITUS ÆTHERIS NITROSI

SPIRIT OF NITROUS ETHER

B P Syn — SWEET SPIRIT OF NITRE

A transparent pale yellow, or greenish-yellow, mobile, volatile and inflammable liquid.

The *B P* describes Spirit of Nitrous Ether as an alcoholic solution containing Ethyl Nitrite, Aldehyde, and other substances; the *U S P* describes it as an alcoholic solution of Ethyl Nitrite (C_2H_5NO), yielding when freshly prepared not less than 4 p.c. of Ethyl Nitrite, the *P G* gives no description.

It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere, and should be exposed as seldom as possible to contact with the air and light.

Medicinal Properties — Stimulant, diaphoretic, diuretic, and antipyretic. Useful in dropsy of renal origin, but is contra-indicated in acute nephritis. Being a nitrite, it is sometimes used in asthma, angina pectoris, and dysmenorrhœa. See also Medicinal Properties of Liquor Ammonii Acetatis.

Dose — 20 to 40 minims = 1.2 to 2.4 c.c., for repeated administration, for a single administration, 60 to 90 minims = 3.6 to 5.4 c.c.

Incompatibles — Potassium Iodide, Ferrous Sulphate, Tincture of Guaricum, Gallic and Tannic Acids, Antipyrine and Salicylates.

Prescribing Notes — When prescribed with Potassium Iodide, separation of Iodine may be prevented by previously neutralising the free acid in Spiritus Ætheris Nitrosi with Potassium or Sodium Bicarbonate, or the Carbonates. The

incompatibility of Antepyrine and Spiritus Ætheris Nitrosi may be overcome by prescribing them in alkaline solution

The measure of gas evolved on the addition of Potassium Iodide solution is a measure of the acidity of the Spiritus Ætheris Nitrosi under examination. It should not amount to much more than a third of the total gas volume registered

Foreign Pharmacopœias — Official in Belg (*Æther Nitricus Alcoholicus*), sp gr 0.84 to 0.86, Dutch (*Nitriis Æthylicus cum Spiritu*), sp gr 0.84 to 0.85, *Spiritus Ætheris Nitrosi*—Ger, sp gr 0.84 to 0.85, Jap, sp gr 0.84 to 0.85, Russ, sp gr 0.84 to 0.85, Swiss, sp gr 0.845 to 0.855, U.S., sp gr about 0.823 at 25° C (77° F) Ital (*Etere Nitroso Ufficinale*), sp gr 0.85, Mex (*Eter Nitroso Alcoholicado*), Norw (*Æther Nitrosus Spirituosus*), sp gr 0.84 to 0.85, Port (*Acido Azotico Alcoollizado*), Span (*Espiritu de Nitro Dulce*)

Under the name of '*Nitrosyl*' a concentrated form of Nitrous Ether has been introduced, 1 fl oz of which mixed with 19 fl oz of Alcohol (90 p.c.) is stated to be equivalent to *Spiritus Ætheris Nitrosi*

Tests—Spirit of Nitrous Ether has a sp gr of about 0.840. The *B.P.* says 0.823 to 0.842, the *U.S.P.* about 0.823 at 25° C (77° F), the *P.G.* 0.840 to 0.850. When freshly prepared it is neutral in reaction towards Litmus paper, but on keeping it gradually develops acidity, and then has an acid reaction towards blue Litmus paper. When carefully poured upon Ferrous Sulphate Solution acidified with Sulphuric Acid a dark brown or blackish-brown coloration is developed at the junction of the two fluids. The *U.S.P.* states that, if a test-tube be half filled with the spirit and put into a water-bath heated to 65° C (149° F) until it has acquired this temperature, the spirit should boil distinctly upon the addition of a few small pieces of broken glass. It is officially required to contain, when freshly prepared, 2½ p.c. w/w of Ethyl Nitrite, and even when it has been kept for some time, and the vessel which contains it has been occasionally opened, it should yield 2 p.c. w/w of Ethyl Nitrite, or a minimum of 1½ p.c. as gasometrically determined by measuring the volume of Nitric Oxide gas evolved on treating it with Potassium Iodide Solution and Diluted Sulphuric Acid, as described below under the heading of Gasometric Determination. The *U.S.P.* requires that it shall contain not less than 4 p.c. of Ethyl Nitrite as gasometrically determined by the process given in small type under the heading of Gasometric Determination. The *P.G.* does not include a method of assay. Allen's method consists in treating the sample with an acidulated solution of Potassium Iodide and measuring the Nitric Oxide liberated. A nitrometer is filled with strong brine, 5 c.c. of Spirit of Nitrous Ether is introduced, followed by 5 c.c. of a strong Potassium Iodide solution, and then by 5 c.c. of diluted Sulphuric Acid. The nitrometer is agitated busily at intervals, after 5 minutes the liquid is adjusted to the same level, the volume of gas is read off. To calculate the percentage of real Ethyl Nitrite the following data is required —

1. The sp gr of the sample to be examined
2. 23.55 c.c. of Nitric Oxide, measured at ordinary pressure and temperature, weigh 0.03 gramme
3. 30 parts by weight of Nitric Oxide are equivalent to 75 parts by weight of Ethyl Nitrite

The measure of gas evolved on the addition of Potassium Iodide is a measure of the acidity of the Spiritus Ætherei Nitrosi under examination. It should not amount to much more than a third of the total gas volume registered. The following process was suggested (*A J P* '98, 273) for the assay of this preparation. Into a 100 c c flask provided with a loosely fitting stopper place successively 10 c c of Distilled Water, 5 c c of a cold saturated aqueous Potassium Chlorate Solution, 5 c c of the sample to be tested, and 5 c c of a 10 p c Nitric Acid Solution. Insert the stopper and shake frequently for 30 minutes, then add 10 c c of Tenth normal Volumetric Silver Nitrate Solution and shake briskly for 1 minute, add 10 drops of Ferric Ammonium Sulphate Solution, and titrate the excess of Tenth normal Volumetric Silver Nitrate Solution with Tenth normal Volumetric Potassium Sulphocyanate Solution. Each c c of Tenth normal Volumetric Silver Nitrate Solution consumed corresponds to 0.0225 gramme of Ethyl Nitrite. This process was claimed to give higher and more correct results than Allen's nitrometer process, but the nitrometer process was never put forward as an absolutely true one, but as one by which Ethyl and other Nitrites might be estimated with approximate accuracy, and it has fulfilled its expectations admirably.

The more generally occurring impurities are free acid and Aldehyde. The *BP* requires that, when shaken with Sodium Carbonate, no effervescence, or only a very feeble effervescence, should occur. The *USP* requires that it should not effervesce when a crystal of Potassium Bicarbonate is dropped into it. The *PG* fixes a limit of acidity, requiring that 10 c c shall not possess an acid reaction after the addition of 0.2 c c of Normal Potassium Hydroxide, but does not state to what indicator of neutrality, presumably towards Litmus. As regards tests for Aldehyde, the *BP* requires that, when a measured quantity of 10 c c of the spirit is mixed with 5 c c of Volumetric Sodium Hydroxide Solution and 5 c c of Water, it should assume a yellow colour, which should not become brown on standing 12 hours. The *USP* includes a somewhat similar test though less stringent, requiring that the mixture should not turn a decided brown within 12 hours, the *PG* does not include a test for Aldehyde.

Gasometric Determination—If 1 volume of the spirit is agitated briskly at intervals during 5 minutes in a nitrometer filled with saturated brine solution, with 1 volume of Potassium Iodide Solution and 1 volume of Diluted Sulphuric Acid, it should, when recently made yield at the normal temperature [15.5°C (60°F)], and pressure (30 in or 760 mm of Mercury), at least $6\frac{1}{2}$, but not more than 7, volumes of Nitric Oxide gas, *BP*.

A quantity of about 30 grammes of the spirit (which has been previously shaken with 0.5 gramme Potassium Bicarbonate is transferred to a graduated measuring flask of 100 c c capacity and its weight accurately determined. It is then diluted with sufficient Alcohol (94 p c) to produce 100 c c and thoroughly mixed. A measured quantity of 10 c c of this alcoholic solution is introduced into a nitrometer filled with a saturated brine. 10 c c of Potassium Iodide Solution is then introduced, and this in turn followed by 10 c c of Normal Volumetric Sulphuric Acid. The volume of gas evolved is read off when the volume of gas has become constant, usually within 30 to 60 minutes. The number of c c of gas is multiplied by 0.307, and the product divided by one-tenth the original weight of the Spirit of Nitrous Ether taken. At the standard temperature and pressure the quotient will represent the percentage of Ethyl Nitrite in the liquid.

The temperature correction is $\frac{1}{3}$ of 1 p c of the total percentage found, for each one degree, additive if the temperature is below, subtractive if above 25° C (77° F). The barometric correction is $\frac{1}{30}$ of 1 p c for each mm, additive if above, subtractive if below 760, *U S P*

LIQUOR ETHYL NITRITIS SOLUTION OF ETHYL NITRITE

A transparent, colourless, or pale yellow, mobile, volatile, inflammable liquid. It consists of a mixture of 95 parts by volume of Absolute Alcohol with 5 parts by volume of Glycerin, and contains, when freshly made, 3 p c by weight, and even after keeping for some time not less than $2\frac{1}{2}$ p c by weight of Ethyl Nitrite.

It should be kept in well-stoppered small glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from exposure to air and light.

The reasons for its introduction will be found, *P J* (3) LVIII 861

Medicinal Properties.—Similar to those of the other more slowly acting Nitrites.

Dose—20 to 60 minims = 1 2 to 3 6 c c

Experiments testing the physiological activity of the *B P* preparation compared it with a 2 per cent solution of the pure Ethyl Nitrite showed that both were equally active.

Tests.—Solution of Ethyl Nitrite has a sp gr of about 0 825, the *B P* states 0 823 to 0 826. It is not official in either the *U S P* or the *P G*. When carefully poured on to the surface of a cold Solution of Ferrous Sulphate acidified with Sulphuric Acid, a dark brown or blackish-brown ring is developed at the junction of the two liquids. It is officially required to yield, when freshly prepared, at least 7 6 volumes of Nitric Oxide gas, and a solution which has been kept some time, and the vessel containing it having been opened, is required to possess at least five-sixths of the strength indicated, as gasometrically determined by shaking for 5 minutes in a brine-charged nitrometer, 1 volume of the solution with 1 volume of Potassium Iodide Solution and 1 volume of Diluted *S*.

The more generally occurring impurities are free acid and Aldehyde. The solution should not effervesce when shaken carefully with Sodium Bicarbonate, indicating the absence of free acid. A measured quantity of 10 c c, when mixed with an equal volume of a mixture of equal parts of Volumetric Sodium Hydroxide Solution and Water, should not assume a yellow colour, indicating the absence of Aldehydes.

Not Official

SPIRITUS FRUMENTI

WHISKY

The term Whisky is here intended to apply to an alcoholic liquid obtained from fermented Grain by distillation and the product of the Pot Still. Whisky is described in the *U S P* as an alcoholic liquid obtained by the distillation of the fermented mash of Grain, such as Indian Corn, Rye, Wheat and Barley, or other mixtures, but no mention is made as to whether it should be the product of the Pot or Patent Still, it is, however, required to be at least 4 years old. Allen

states in the majority of cases a judicious admixture of raw and malted Grain is employed, other things being equal, the spirit from malted Grain is the most valuable and contains least Fusel Oil. It has recently been held as the result of a magisterial decision that only Pot Still Whisky distilled in Ireland and Scotland could be legally sold as Whisky, thus eliminating Patent Still Whisky. An appeal lodged against this decision resulted in an equal division of opinion on the subject, and the case was referred for re-hearing. The Royal Commission appointed to investigate not only the point whether Pot or Patent Still Whisky should be the legal representative of the article, but also the whole position with regard to the subject, in an interim report arrives at the following conclusions:— (1) That no restrictions should be placed upon the processes of, or apparatus used for the distillation of any Spirit to which the term Whisky may be applied as a trade description. (2) That the term Whisky having been recognised in the past as applicable to a potable Spirit manufactured from (i) Malt, or (ii) Malt and other malted Barley, or other cereals, the application of the term Whisky should not be denied to the product manufactured from such materials. Whisky improves greatly on keeping, when new it is colourless or nearly so, but by storing in Sherry casks (a favourite method of imparting flavour to Whisky) it acquires colour, and then contains sensible traces of Tannin, Sugar, etc. The smoky flavour of Irish Whisky is due to the fact that the Malt used has been dried upon kilns in which Peat is used for fuel, but is sometimes imitated by adding 1 or 2 drops of Cresosote to the gallon of spirit. It is doubtful whether Fusel Oil is ever purposely added to Whisky, but it is almost invariably present in greater or less quantity, and has been stated to be the cause of objectionable symptoms produced by new spirit. Allen is of opinion that as the Amyl Alcohol in spirits rarely exceeds 0.1 p.c. or 70 grains per proof gallon, it seems highly improbable that it could produce the local effects sometimes attributed to it, its effect on the general system has probably been greatly exaggerated. It is a noticeable fact (*B M J* '03, ii 1645) that whereas years ago 70 p.c. of the Whisky was Malt Whisky and 30 p.c. Grain or Patent Spirit, the proportions are now reversed.

Tests—Whisky has a sp. gr. of about 0.930. The *U.S.P.* states it should not be more than 0.945 nor less than 0.924 at 15.6° C (60° F).

Absolute Alcohol—It contains from 50 to 64 p.c. w/v of Absolute Alcohol. The *U.S.P.* says from 44 to 55 p.c. w/v corresponding to 37 to 47.5 p.c. by weight of Absolute Alcohol. The Alcohol may be determined by distilling a known volume of the Whisky and ascertaining the sp. gr. of the distillate when made up to a definite volume. The percentage of Absolute Alcohol by weight, corresponding to this gravity, may be found by reference to an Alcohol table. In calculating the result on the original Whisky it is necessary to take into consideration the sp. gr. of the sample and the volume used for the distillation.

Extractive Matter—The amount of extractive matter rarely amounts to more than 100 grains per gallon, equivalent to about 0.15 p.c. The *U.S.P.* states that the limit of residue when dried at 100° C (212° F) is 0.5 p.c. w/v, and this residue is required to possess no sweet or distinctly spicy taste. The amount may be determined by evaporating a definite volume (25 c.c.) to dryness on a water bath, drying the residue at a temperature of 100° C (212° F) till constant in weight, and when cool, weighing.

Total Acid—The total acid may vary from 0.01 p.c. to 0.083 p.c., and may be determined by titrating a measured quantity of 25 c.c. of the sample with Tenth normal Barium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. The *U.S.P.* requires that to render 100 c.c. of Whisky distinctly alkaline to Litmus not more than 1.2 c.c. of Normal Volumetric Potassium Hydroxide Solution should be necessary.

Volatile Acid—The volatile acid may vary from 0.014 p.c. to 0.072 p.c. w/v. It may be determined by distilling a measured quantity (100 c.c.) almost to dryness, adding 25 c.c. of Water to the residue in the distillation flask and continuing the distillation until reduced to a low bulk. The distillate is titrated with Tenth-normal Volumetric Barium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 c.c. of Tenth normal

Volumetric Barium or Sodium Hydroxide Solution is equivalent to 0.005958 gramme of Acetic Acid. The difference between the number of c.c. of Tenth-normal Volumetric Barium or Sodium Hydroxide Solution required to neutralise the volatile acid, and the calculated number of c.c. of Tenth-normal Volumetric Barium or Sodium Hydroxide Solution required to neutralise 100 c.c. of the original sample represents that necessary for the neutralisation of the fixed acid, and may be calculated into Tartaric Acid. Each c.c. of Tenth-normal Volumetric Barium or Sodium Hydroxide Solution represents 0.007446 gramme of Tartaric Acid.

Esters—The proportion of esters in Whisky varies from 0.033 p.c. w/v to 0.185 p.c. w/v. They may be determined by distilling a measured quantity of 100 c.c. of the sample in a distillation flask almost to dryness, adding 25 c.c. of Water and again distilling to a volume of about 5 c.c. The free acid in the distillate is neutralised with Tenth-normal Volumetric Barium or Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. A measured quantity of 25 c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution is then added and the contents of the flask boiled under a reflux condenser for 1 hour, cooled, and the excess of Tenth-normal Volumetric Alkali Solution titrated with Tenth-normal Volumetric Sulphuric Acid Solution, still employing Phenolphthalein Solution as an indicator of neutrality. In the event of the specimen containing more Ethers than correspond with this quantity of Sodium Hydroxide Solution, which is at once indicated by the disappearance of the pink colour, a further quantity of the Tenth-normal Volumetric Alkali Solution should be added. The number of c.c. of Tenth-normal Volumetric Sulphuric Acid Solution used is subtracted from the number of c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution added, the difference is calculated into terms of Ethyl Acetate. 1 c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution corresponds to 0.0088 gramme of Ethyl Acetate.

Higher Alcohols—The higher Alcohols in Whisky vary from 0.082 p.c. w/v to 0.260 p.c. w/v. Various methods have been suggested for the determination of higher Alcohols, the most reliable and the one now most generally used is the Allen-Marquardt process as modified by Allen and Chattaway, and generally referred to as the Allen-Marquardt process. It consists in distilling a measured quantity of 200 c.c. of the specimen to be examined until reduced in volume to 20 c.c. To the residue in the distillation flask are added 50 c.c. of Water and 10 grammes of Sodium Sulphate, and the distillation continued until reduced to 10 c.c., the mixed distillates are titrated with Tenth-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. A further measured quantity of 20 c.c. of Tenth-normal Volumetric Sodium Hydroxide Solution is added and the liquid boiled for 1 hour under a reflux condenser, the excess of Tenth-normal Volumetric Sodium Hydroxide Solution is titrated with Tenth-normal Volumetric Hydrochloric Acid Solution. The liquid is now divided into two equal parts, each corresponding to 100 c.c. of the original

which is treated in the following way in order to yield a
A saturated solution of common salt (prepared by
lean table salt and adding dilute Sulphuric Acid until
the liquid has a distinct acid reaction, and filtering) is added to the liquid until
the resulting mixture has a sp. gr. of 1.1 when it is extracted in a separator
four times with Carbon Tetrachloride, using 40 c.c. of the Carbon Tetrachloride
for the first extraction, 30 c.c. for the second, 20 c.c. for the third, and 10 c.c. for
the last extraction. The Carbon Tetrachloride now contains the whole of higher
Alcohols and some Ethyl Alcohol. To remove the latter the Carbon Tetra-
chloride is shaken with 50 c.c. of brine, and after this has been separated it is
shaken with 50 c.c. of a saturated Solution of Sodium Sulphate to remove the
Chloride. The Carbon Tetrachloride is next treated with an oxidising mixture
consisting of 5 grammes of Potassium Bichromate, 2 grammes of strong Sulphuric
Acid and 10 c.c. of Water, the oxidation may be carried out in a flask connected
with a reflux condenser, or preferably in well-made glass bottles possessing
accurately fitting ground glass stoppers, the stoppers being securely tied down
and the bottles heated in a water-oven. When the oxidation is performed
under a reflux condenser, at least 8 hours boiling is necessary, but when
conducted under pressure in stoppered bottles, if shaken frequently, from 3 to 4

hours only is necessary for complete oxidation. The liquid is transferred to a distilling flask, the bottle rinsed with 30 c.c. of Water, the washings transferred to the distilling flask and the liquid distilled until only 20 c.c. remain in the flask. A measured quantity of 80 c.c. of Water is added to the residue and the distillation continued until only 5 c.c. remain. The mixed distillates are now titrated with Tenth normal Volumetric Barium Hydroxide Solution using Methyl Orange Solution as an indicator of neutrality, shaking the liquid thoroughly after each addition. The amount of alkali required to neutralise the liquid at this stage should not exceed 2 c.c., and generally less is required, a few drops of Phenolphthalein Solution are now added and the titration continued. Each c.c. of Volumetric Barium Hydroxide Solution required in the second stage of the titration corresponds to 0.0089 gramme of higher Alcohols expressed in terms of Amyl Alcohol. The alkali which was added when titrating with Methyl Orange Solution is stated to represent the mineral acid which is distilled. This method has been carefully investigated by Schidrowitz and Kaye (*J S C I* '02, 815, *Analyst* xxx 190), who state that 'of the methods examined, that of Allen Marquardt, with certain minor modifications, is alone capable of giving fully reliable figures, at any rate where Whisky is concerned, and that we are still inclined to believe that if carefully worked this process is still the most reliable of all those published'. The process has also been criticised (*J S C I* '06, 1125), in the course of an extended investigation of samples of Whisky for the Western Australian Government it was found that operating upon pure materials no mineral acid was produced as the result of the oxidising mixture alone or from its action upon Carbon Tetrachloride. It was, however, noticed that Valerianic Acid in a similar manner to Acetic Acid had a distinct action upon Methyl Orange Solution, and it is concluded that the titration for mineral acids is unnecessary and introduces errors, and that accurate results may be obtained by observing the following points: (a) The shaking out should be performed at a temperature of 15.5° C (60° F) or less. (b) the oxidation to be conducted in pressure bottles, (c) the higher Alcohols should be determined by direct titration only, calculating all acidity as Valerianic Acid.

Aldehyde—Aldehyde may be detected in the distilled spirit, or if present in very minute proportions in the first fraction of the distillate. 10 c.c. of the distilled spirit may be mixed with 4 c.c. of Schiff's reagent, prepared by mixing 30 c.c. of a 1 in 1000 aqueous solution of Magenta with 20 c.c. of Sodium Bisulphite Solution (sp gr 1.81), 3 c.c. of Sulphuric Acid and 200 c.c. of Water.

Furfural—Furfural may be detected by means of Aniline Acetate, 10 c.c. of the distilled spirit is mixed with 2 c.c. of an Aniline Acetate Solution prepared by dissolving 10 drops of Aniline in 2 c.c. Glacial Acetic Acid.

As a test for the absence of more than a trace of Fusel Oil from grain, the *U S P* requires that if 100 c.c. of Whisky be very slowly evaporated on a dish or water bath, the last portions volatilised should not have a harsh or disagreeable odour. The absence of added Sugar, Glycerin and aromatic substances, is judged by the character of the residue left on drying as described above. The absence of more than traces of Oak Tannin from casks is assured by the residue being required to dissolve completely in 10 c.c. of cold Water to form a solution which should not be coloured deeper than light green on the addition of a few drops of Ferric Chloride T.S. The *U S P* also requires that if 50 c.c. of Whisky be shaken vigorously in a stoppered flask with 20 c.c. of Kaolin, and filtered, after standing half an hour the filtrate should not be lighter in colour than the Whisky before treatment.

Official in U S

Not Official

SPIRITUS METHYLATUS

METHYLATED SPIRIT

The duty free spirit supplied to 'manufacturers' under a special bond, is a mixture of 9 parts of Alcohol with 1 part of a Wood Naphtha, approved by the Excise. It can also be supplied under a special bond for scientific purposes.

As supplied to 'licensed retailers,' Methylated Spirit is 3 pints of Petroleum Oil added to 100 gallons of the mixture described above. The Petroleum Oil is added, partly to make it more nauseous for drinking, and partly to facilitate its recognition. It becomes turbid when mixed with Water, which quality renders it unsuitable for many purposes to which duty-free spirit has been applied.

Licensed retailers of Methylated Spirit must not sell more than 1 gallon at any one time, and may not keep stock exceeding 50 gallons. They may not sell Methylated Spirit between the hours of 10 p.m. on Saturdays and 8 a.m. on Mondays.

SPIRITUS RECTIFICATUS.

ALCOHOL (90 p.c.)

B.P. Syn—RECTIFIED SPIRIT

A transparent, colourless, mobile, volatile and inflammable liquid, having a distinctive spirituous odour and burning taste.

The Alcohol (90 p.c.) of the *B.P.* is described as a liquid consisting of 90 parts by volume, equivalent to 85.65 p.c. by weight, of Ethyl Hydroxide, C_2H_5OH , eq. 45.7, and 10 parts by volume, equivalent to 14.35 p.c. by weight, of Water, and its official method of preparation is by the distillation of fermented Saccharine liquids. It is only slightly stronger than the Rectified Spirit of the *B.P.* '85, containing, by volume, 1.35 p.c., or by weight 1.65 p.c. more Ethyl Hydroxide. The equivalent to this spirit in the *U.S.P.* is known under the title Alcohol, and it is described as a liquid composed of about 92.3 p.c. by weight, or about 94.9 p.c. w/v of absolute Ethyl Alcohol, and about 7.7 p.c. w/w of Water. The *P.G.* describes it under the heading of Spiritus, and states that 100 parts contain 91.2 to 90 parts by volume, equivalent to 87.2 to 85.6 p.c. by weight.

It should be kept in well closed vessels and in a cool atmosphere, and it should be kept away from lights or fire.

On mixing Alcohol (90 p.c.) and Water, contraction of volume and rise of temperature occur. When such a mixture is prescribed in the British Pharmacopœia, the cooled liquid should be employed.

It is possible to rectify Alcohol up to 98 p.c., and 95 p.c. is prepared commercially in large quantities.

It may here be noted that although it is illegal for chemists and druggists to sell Rectified Alcohol except upon prescription, the Board of Inland Revenue do not intend to interfere with its sale by them in small quantities not exceeding 8 oz. or a lb., for the purposes of medical or scientific research.

Alcohol (90 p.c.) dissolves Camphor, Balsams, Castor Oil, Iodine, Potassium and Sodium Hydroxides, but not the Carbonates.

Medicinal Properties—Internally a powerful diffusible stimulant, especially cardiac, antipyretic, diuretic, and diaphoretic. Used in some states of acute disease characterised by excessive debility, as in typhoid, acute pneumonia, and influenza, to maintain the strength over the crisis, in chronic wasting diseases as phthisis, in insomnia of old people, during a meal in small quantity, as an aid to digestion and absorption, and to promote appetite, more especially in the aged and feeble and in those exhausted by overwork, in sudden fainting. In acute dyspepsia it is injurious, it may, however, check vomiting, and brandy often checks diarrhoea. In moderation

it acts as a food, and saves tissue waste. Large quantities paralyse the gastric secretion, produce gastritis, and ultimately atrophy of the gastric glands. Externally to prevent bed sores and cracked nipples by hardening and disinfecting the skin, it is antiseptic and astringent, and is applied diluted to stop sweating and to produce cold by evaporation, 1 of Alcohol (90 p c) and 2 of Camphor Water mixed is a good evaporating lotion. Diluted it forms a lotion for eczypelas, erythema, burns, and scalds while the cuticle is entire, and for sprains and recent bruises. As an ingredient of liniments it is rubefacient, it relieves rheumatic and other kinds of pain, and aids the resorption of inflammatory products.

Foreign Pharmacopœias—Official in all Fr., Alcool éthylique à 95 degrés centésimaux.

Tests—Alcohol (90 p c) has a sp gr of 0.834. The sp gr of the Alcohol official in the *USP* is about 0.816 at 15.6° C (60° F) or 0.809 at 25° C (77° F), that official in the *PG* 0.830 to 0.834. It readily volatilises, and when ignited burns with a pale blue non-luminous flame. If to a few c c of Alcohol 1 or 2 drops of a Potassium Hydroxide Solution (50 p c) be added, and then a solution of Iodine drop by drop till the liquid contains a slight but distinct excess of Iodine, and the mixture be warmed to about 60° C (140° F), it yields a powerful penetrating odour of Iodoform. When mixed with a small quantity of Solution of Potassium Permanganate and Diluted Sulphuric Acid it is rapidly oxidised, the distinctive odour of Acetaldehyde being evolved. It reduces Potassium Bichromate Solution, yielding similarly an odour of Acetaldehyde, the solution changing to a bright green. It should be miscible in all proportions with Water, the solution should be odourless and free from turbidity.

The more generally occurring impurities are fixed residue, oily or resinous substances, Fusel Oil and allied impurities, Amylic Alcohol and other organic impurities, Aldehyde, Tannic Acid from Oak cask, excess of Aldehyde. It leaves no weighable residue upon evaporation, indicating the absence of fixed residue. It should afford a clear liquid when mixed with Water, indicating the absence of oily or resinous substances. A small quantity of the Alcohol allowed to evaporate on a piece of clean white bibulous paper should leave no unpleasant odour after the Alcohol has completely evaporated, indicating the absence of Fusel Oil and allied impurities. The *USP* mixes the Alcohol with half its volume of Water and one-tenth of its volume of Glycerin and allows the mixture to evaporate spontaneously. The *PG* does not include an evaporation test. The remarks upon the *BP* and the *USP* methods of testing for Fusel Oil and allied impurities are described under Alcohol Absolutus. The *PG* requires that, when a mixture of 10 c c of spirit and 0.2 c c of Potassium Hydroxide Solution (15 p c) is evaporated to one-tenth its volume the residue, when supersaturated with Sulphuric Acid, should develop no odour of Fusel Oil. As an additional test for the presence of Amylic Alcohol and readily charred organic impurities, the *USP* evaporates 25 c c spontaneously in a porcelain evaporating dish,

carefully protecting the liquid from dust during the evaporation. The evaporation is continued until the surface of the dish is barely moist, on then adding a few drops of colourless concentrated Sulphuric Acid the residue should not produce a red or brown coloration. The *P G* performs the test for readily carbonisable organic impurities on the spirit direct without evaporation. In testing for Aldehyde the *BP* and *USP* give practically the same test. The Ammonia test for Tannic Acid and excess of Aldehyde is common to the *BP* and the *P G*, but is omitted from the *USP*. The *P G* includes, in addition, a test with Hydrogen Sulphide which is described below. The *USP* includes a test for the absence of more than 2 p.c. of Methyl Alcohol. The explanation of this test, given under Alcohol Absolutum and a description of the test itself, is in small type below, it is not given in either the *BP* or the *P G*.

Potassium Hydroxide—On mixing 10 c.c. of the spirit with 5 c.c. of Potassium Hydroxide TS the liquid should not immediately darken in colour, *BP*, should not at once assume a yellow colour, *USP*.

Ammonia—No immediate darkening in colour should occur on the addition of Ammonia TS to Alcohol (90 p.c.), *BP*, the spirit should not become coloured, *P G*.

Hydrogen Sulphide—Spirit should not become coloured by TS of Hydrogen Sulphide, *P G*.

Silver Nitrate—After exposing 100 c.c. of Alcohol (90 p.c.) with 2 c.c. of Volumetric Silver Nitrate Solution to bright light for 24 hours and decanting the liquid from the black powder formed, no further change should occur when the liquid is again exposed with more of the Volumetric Silver Nitrate Solution, *BP*, 10 c.c. of spirit should neither become turbid nor coloured on warming with 5 drops of TS of Silver Nitrate, *P G*, if 20 c.c. of Alcohol be shaken in a clean, vial with 1 c.c. of Silver Nitrate TS, the mixture should not than faintly opalescent or acquire more than a faint brownish tint when exposed for 6 hours to diffused daylight, *USP*.

Sulphuric Acid—A mixture of 10 c.c. of spirit and 0.2 c.c. of Potassium Hydroxide, evaporated to 1 c.c. and supersaturated with diluted Sulphuric Acid, should not have any odour of Fusel Oil, *P G*. If 5 c.c. of spirit be carefully poured as a layer over 5 c.c. of Sulphuric Acid, no rose red zone should form at the line of contact, even after standing for some time, *P G*. If 25 c.c. of Alcohol be allowed to evaporate spontaneously in a porcelain evaporating dish, carefully protected from dust, until the surface of the dish is barely moist, no red or brown colour should be produced upon the addition of a few drops of colourless, concentrated Sulphuric Acid (absence of Amyl Alcohol, or non-volatile, carbonisable, organic impurities), *USP*.

Potassium Permanganate—The red colour of a mixture of 10 c.c. of spirit and 1 c.c. of Potassium Permanganate TS should not become yellow within 20 minutes, *P G*.

Copper Wire and Resorcin—1 c.c. of the Alcohol or spirit is transferred to a test-tube of a capacity of about 40 c.c., and sufficient Water added to bring the volume of the liquid to 10 c.c., the test necessitating the use of 1 c.c. of Al in the liquid not being more than about 10 p.c. by volume. 1 metre of No. 18 clean Copper Wire is wound closely round a glass rod 7 mm. thick so as to form a coil about 3 cm. long, the end of the wire being formed into a handle, the coil is heated to redness in a non-luminous flame and plunged, whilst red hot, to the bottom of the liquid in the test-tube and held there for a second or two, withdrawn, and dipped into Water to cool. This operation is repeated five or six times, immersing the test-tube in cold Water to keep it cool. The contents are now filtered into a wide test-tube and boiled very gently, if the odour of Acetalde-

hyde be perceptible the boiling is continued until the odour has ceased to be clearly distinct, the liquid is cooled, a drop of a 1 in 200 Resorcinol solution added. A portion of this liquid is poured carefully upon the surface of some pure concentrated Sulphuric Acid contained in another test tube, the tube is allowed to stand for 3 minutes and then slowly rotated, a rose red coloration should not develop at the point of contact of the two liquids (absence of more than 2 p c of Methyl Alcohol), *U S P*

DILUTED ALCOHOL

Four strengths of diluted Alcohol are official containing respectively, 70, 60, 45 and 20 p c of Ethyl Hydroxide by volume. They may be prepared as described in the following paragraphs.

1 Alcohol (70 p c)—124½ fl oz of Alcohol (90 p c) mixed with 38½ fl oz of Water, or 777.7 c c of Alcohol (90 p c) with 241.6 c c of Water, temperature 15.5° C (60° F) Sp gr 0.8900

2 Alcohol (60 p c)—With 106½ fl oz of Alcohol (90 p c) mix 57½ fl oz of Water, or with 666.6 c c of Alcohol (90 p c) mix 357.8 c c of Water, temperature 15.5° C (60° F) Sp gr 0.9135

3 Alcohol (45 p c)—With 80½ fl oz of Alcohol (90 p c) mix 84½ fl oz of Water, or with 500 c c of Alcohol (90 p c) mix 526.6 c c of Water, temperature 15.5° C (60° F) Sp gr 0.9436

4 Alcohol (20 p c)—With 35½ fl oz of Alcohol (90 p c) mix 126½ fl oz of Water, or with 222.2 c c of Alcohol (90 p c) mix 791 c c of Water, temperature 15.5° C (60° F) Sp gr 0.9760

When the sp gr of Alcohol is 0.920 it is called **Proof Spirit**, if lighter than this, it is called 'above proof', if heavier than this, 'under proof', and the percentage of Water, or of Rectified Spirit, sp gr 0.825 (the Inland Revenue Standard), by measure, necessary to be added to any sample of Spirit to bring it to the standard of Proof Spirit, indicates the number of degrees the given sample is above or below proof. Thus, if 100 volumes of a Spirit require 10 volumes of Water to reduce it to proof, it is said to be '10 over proof' on the other hand, if 100 volumes of Spirit require 10 volumes of Spirit to raise it to proof, the sample is said to be '10 under proof'.

U S defines three strengths of Alcohol. **Alcohol Absolutum**, containing 99 p c of Alcohol, **Alcohol**, about 94.9 p c, and **Alcohol Dilutum**, about 48.9 p c. All by volume.

Ger describes four strengths. **Alcohol Absolutus**, containing 99.4 to 99.7 p c of Alcohol, **Spiritus**, 90 to 91.2 p c, **Spiritus Dilutus**, 68 to 69 p c, **Spiritus e Vino**, 37 to 41 p c. The three former by volume, the last by weight.

The Spirits of the Pharmacopœias are as follows:—

	Sp gr	Percentage of Absolute Alcohol by Volume	
British	0.834	Alcohol 90 p c (Spiritus Rectificatus)	90
"	0.890	" 70 p c	70
"	0.9135	" 60 p c	60
"	0.9436	" 45 p c	45
"	0.976	" 20 p c	20
Austrian	0.830—0.834	Sp Vini Concentratus	90 to 91.2
"	0.892—0.896	" Dilutus	68 to 69
"	0.935—0.945	" Cognac	44 to 48
Belgian	0.816—0.820	" Spiritus	94.09
Danish	0.812—0.816	Spiritus Alcoholisatus	95 to 96
"	0.831—0.834	" Concentratus	90 to 91
"	0.890—0.895	" Dilutus	68 to 70
Dutch	0.8159	" Fortior	95
"	0.8897	" Dilutus	70
French	0.79433—0.8095	Alcool Ethylique	100
"	0.816	" " at 95°	95

	Sp Gr	Percentage of Absolute Alcohol by Volume
German	0 796—0 800	Alcohol Absolutus 99 4 to 99 7
"	0 830—0 834	Spiritus 90 to 91 2
"	0 892—0 896	" Dilutus 68 to 69
"	0 920—0 924	" o Vmo (by weight) 37 to 41
Hungarian	0 831—0 834	" 90 to 91
"	0 892	" Dilutus 70
"	0 919—0 924	Cognac (by weight) 46 to 50
Italian	0 8346	Alcool 90° 90
"	0 800	" Assoluto 99
"	0 9141	" 60° 60
Japanese	0 830—0 834	Spiritus 90 to 91 2
"	0 892—0 896	" Dilutus 68 to 69
Mexican	0 79	Alcohol Vinico 100
"		" at 50° 50
"		" 60° 60
"		" 80° 80
"		" 90° 90
Norwegian	0 8306—0 8339	Spiritus Concentratus 90 to 91
"	0 9021—0 9044	" Dilutus 64 to 65
Portuguese	0 834	Alcool at 90° 90
"	0 850	" 85° 85
"	0 905	" 65° 65
Russian	0 813—0 816	Sp Vini Alcoholicatus 95
"	0 831—0 834	" Rectificatus 90
"	0 888—0 890	" Rectificatus 70
"	0 952—0 955	" Dilutus 38
Spanish	0 794	Alcohol Anhidro 100
"	0 8161	" de 95° 95
"		" de 60° 60
Swedish	0 831—0 833	Spiritus Concentratus 90 to 91
"	0 903—0 905	" Dilutus 63 to 64
"	0 935	" Tenuis 48 to 50
Swiss	0 830—0 834	" 90 09 to 91 29
"	0 892—0 895	" Dilutus 68 12 to 69 34
"	0 916—0 939	" e Saccharo (Rum) 50 to 60
"	0 927—0 950	" e Vmo (Cognac) 45 to 55
U S	0 816	Alcohol 94 9
"	0 797	" Absolutum (by weight) 99
"	0 936	" Dilutum 48 9
"	0 925—0 941	Sp Vini Gallici 46 to 55
"	0 924—0 945	" Frumenta 44 to 55

Relative Strength of Wines and Spirits—The following figures represent the average strength in Alcohol by Volume Jamaica Rum, about 69 p c , Proof Spirit, about 57 p c , Whisky, about 51 p c , Brandy, about 48 p c , Gin, about 47 p c , Port, Sherry and Madena, about 20 p c , Benedictine, about 15 p c , Hock and Moselle, about 10 p c , strong Ale and Stout, 7 to 8 p c , Beer and Cyder, 5 to 6 p c

SPIRITUS VINI GALlici.

BRANDY

The *B P* describes Brandy as a spirituous liquid distilled from Wine and matured by age. It is required to contain not less than 36½ p c w/w or 43½ p c w/v of Ethyl Hydroxide. The *U S P* describes it as an alcoholic liquid obtained by the distillation of the fermented unmodified juice of fresh grapes, requires it to be at least 4 years old, and that it shall contain approximately 39 to 47 p c w/w or 46 to

55 w/v of Absolute Alcohol. The *P G* does not include Brandy. A good deal of attention has been devoted to the subject of Brandy, and numerous prosecutions have arisen from the sale of a mixture of Brandy and a spirit not derived from the distillation of the grape as genuine Brandy. At a discussion opened before the Society of Public Analysts by Mr Otto Hehner and reported in the *Analyst* xxx 36, it seemed to be generally agreed that a definite standard of 80 per 100,000, that is to say, 80 grammes of Ethyl Acetate per 100,000 of Absolute Alcohol, could not be relied upon for the compound Ethers. Schidrowitz and Kaye have found (*Analyst*, xxx 149) that the 'break down' of Brandy if effected with faintly alkaline Water is liable to seriously affect the Ether value, a sample possessing an Ester value of 98.8, that is to say, containing 98.8 grammes of Ethers expressed as Ethyl Acetate per 100 litres of Absolute Alcohol, when broken down the Ester value was reduced to 66.5. The former author considers (*J S C I*, '05, 177) that each case should be considered on its merits, and no conclusion should be drawn from a single figure, on the other hand the sample which is analytically satisfactory is not necessarily genuine. In cases of doubt a taster's assistance should be requisitioned, if both the opinion of the analyst and of an unprejudiced taster are adverse to the sample, the merchant or distiller should be called on to give evidence as to the origin, etc.

In the present state of our knowledge it is impossible to tell from an analytical point of view the origin of the spirit in Brandy.

Tests—The *B P* does not include any test for Brandy, nor does it even indicate a method by which the official percentage of Ethyl Hydroxide may be ensured. The sp gr of Brandy is about 0.930. The *USP* requires that the sp gr of Brandy should be not more than 0.941 nor less than 0.925 at 15.6° C (60° F). Genuine typical Brandy contains from 39 to 47 p c by weight of Absolute Alcohol, and these limits have been adopted as a standard of strength by the *USP*. The average strength of Absolute Alcohol is about 42 p c w/w. The percentage of Absolute Alcohol may be determined by a similar method to that described under Spiritus Frumenti. The extractive matter varies from 0.6 to 1.5 p c and averages about 0.75 p c w/v. The *USP* requires that the percentage w/v of residue dried at 100° C (212° F) should not amount to more than 0.5, and that this residue should have no sweet or distinctly spicy taste, indicating the absence of added Sugar, Glycerin and aromatic substances, and may be determined by evaporating a measured quantity of 25 c c to dryness on a water-bath, the residue being dried at a temperature of 100° C (212° F) until constant in weight, and when cool, weighed. The proportion of volatile acid varies from 0.032 p c w/v to 0.1 p c w/v, averaging about 0.042 p c w/v. It may be determined on the distilled spirit by a similar method to that described under Spiritus Frumenti, and may be expressed in terms of Acetic Acid. The compound Ethers expressed in terms of Ethyl Acetate vary from 0.051 to 0.086 p c w/v, and average about 0.055 p c w/v, equivalent to 130.9 per 100,000, calculated on a spirit of an average Alcohol content of 42 p c.

The amount may be determined by the Saponification test given under Spiritus Frumenti. The *U S P* includes a limit of free acid which represents the total acidity of the Brandy, volatile acid as well as fixed, and mentions that 100 c c of the Brandy should require not more than 1 c c of Normal Volumetric Potassium Hydroxide Solution to render it distinctly alkaline to Litmus. The higher Alcohols expressed in terms of Amyl Alcohol vary from 0.05 p c w/v to 0.136 p c w/v, averaging about 0.068 p c w/v, then amount may be determined by the Allen-Marquardt process given under Spiritus Frumenti, and the same tests as are there described for the detection of Aldehyde and Furfural may also be employed here.

The Circular No. 18 issued by the United States Department of Agriculture defines potable Brandy as a distillate from Wine properly aged by storage in wood, to eliminate the amount of Fusel Oils, etc., which may be present. It is required to contain not less than 45 nor more than 55 p c w/v of Absolute Alcohol, and not more than 0.25 p c w/v of extractive matter. The contents of Fusel Oil should not exceed 0.25 p c w/v. Brandy should not be mixed with Alcohol from any other source than that of distilled Wine, the distillate from the lees, pomace, and refuse of the winery is not entitled to the term 'Brandy' in the potable sense. Cognac is only a name, in the case of Brandies made in Cognac from wine of at manufacture there, no artificial colour other than that distilled in the wood in which they are aged is admitted in Brandies. It requires that when 100 c c of Brandy are slowly evaporated in a dish on a water-bath the last portions volatilised should be odour free from harshness, indicating the absence of Oil from grain or pot spirit, the residue should completely dissolve in 10 c c of cold Water, and the solution so produced should not be coloured deeper than light green on the addition of a few drops of Ferric Chloride T.S., indicating the absence of more than traces of Oak Tannin from casks.

Foreign Pharmacopœias—Official in Austria (Spiritus Vini Cognac), U.S. (Spiritus Vini Gallici).

Preparation

MISTURA SPIRITUS VINI GALLICI. MIXTURE OF BRANDY

Rub the yolks of two Eggs with $\frac{1}{2}$ oz of Refined Sugar, add 4 fl oz of Cinnamon Water and 4 fl oz Brandy.

Dose.—As a draught, 1 to 2 fl oz = 28.4 to 56.8 c c

Not Official

STANNI OLEAS

A greyish, coarsely granular powder, insoluble in Alcohol, very slightly soluble in Almond Oil, completely disintegrated and partially dissolved by Ether or Oleic Acid.

UNGUENTUM STANNI OLEATIS—Stannous Oleate, 60 grains, Lard, 1 oz.

Of great utility in diseases of the nails, it overcomes the brittle, split and soft conditions of the nails, and gives them a brilliant lustre.—*B.M.J.* '84, ii, 753, *T.G.* '86, 494.

STAPHISAGRIÆ SEMINA.

STAVESACRE SEEDS

FR, STAPHISAICRE, GER, SILPHANSKORNEL, ITAL, STAFISAGRIA,
SIAM, ES-LATISACRIA

The dried ripe Seeds of *Delphinium Staphisagria*

Medicinal Properties — The Seeds have been used in ointments for many years as a parasiticide for pediculi, the activity rests in an Oil which they contain in rather large quantity. The late Balmanno Squire experimented with this Oil, and also with the Seeds from which the Oil had been withdrawn by Ether, and found the latter inert. He successfully used an ointment made with the Oil in prurigo semilis.

Official Preparation — Unguentum Staphisagrie

Not Official — Lotio Staphisagrie, Oleum Staphisagrie, Unguentum Olei Staphisagrie, and Delphinina

Foreign Pharmacopœias — Official in Belg, Dutch, Fr, Ital, Port, Mex and U S

Descriptive Notes — The Seeds are greenish black, or blackish-brown if not quite ripe, about three lines long, and rather less in width, irregularly 4 to 5 sided with one side convex and the others more or less flattened or concave, the angles are sharp, and the testa is rough, wrinkled and deeply pitted. The albumen is whitish, oily, and has a minute embryo at the pointed end. No other *Delphinium* in cultivation has so large a seed, the species usually grown in botanic gardens as this plant has smaller seeds and lilac flowers and is *D. pictum*, Willd. *D. Staphisagria*, L., has blue flowers and is only half hardy.

Tests — Stavesacre Seeds yield from 10 to 15 p c of ash. The Seeds contain a large proportion of fixed Oil. Four samples of Seeds examined in the author's laboratory when extracted with Ether yielded 31.4, 32.8, 33.9 and 34.8 p c of Oil.

Preparation

UNGUENTUM STAPHISAGRIÆ STAVESACRE OINTMENT

Digest 2 of crushed Stavesacre Seeds in 8½ of Benzoated Lard on a water-bath for 2 hours, squeeze through calico, and dissolve 1 of Yellow Beeswax in the hot liquid, and finally strain until cold.

About half the strength of B P '85, and Yellow Beeswax is added.

Official in Ital, 1 and 3

Not Official

LOTIO STAPHISAGRIÆ *Syn.* Nuisery Hair Lotion — Stavesacre Seeds, in rough powder, 2 oz, Acetic Acid, 1 oz, Water, 16. Boil for 10 minutes in a covered vessel, set aside till cold, then add Rectified Spirit, 2 oz, Oil of Geranium, 2 minims, Oil of Lavender, 2 minims, Oil of Lemon, 4 minims, filter and add Glycerin, 1 oz, Water, to 20 fl oz. This is the Edinburgh Infirmary Pharmacopœia preparation — *Pharm. Form.*

This has been incorporated in the B P C

OLEUM STAPHISAGRIÆ—The Oil obtained by expression from the Seeds

It is insoluble in Alcohol (90 p c), but dissolves readily in hot Absolute Alcohol

Tests—Stavesac Oil has a sp gr of about 0.918

UNGUENTUM OLEI STAPHISAGRIÆ—Expressed Oil, 60 minims, Lard, 1 oz Used as a non-irritant remedy in scabies and in phtheiasis

DELPHININA Delphinine—Rhombic crystals or as a yellowish powder An alkaloid obtained from Stavesac Insoluble in Water, but dissolves in acidulated Water, in Alcohol, Ether and Chloroform It melts at about 192° C (377.6° F) It yields no colour reactions with acids, but when mixed with 1 to 2 volumes of Amyl Alcohol and treated with Sulphuric Acid, it yields an orange-coloured mass, which after several hours becomes dark rose-red and ultimately blue

Dose— $\frac{1}{10}$ grain = 0.0011 gramme, and repeat every 2 hours in neuralgia
—*L. M. R.* '87, 446, *J.* '87, n. 879

Not Official STEARIN.

COCOA-NUT STEARIN

A white, soft, crystalline, fatty substance, unctuous to the touch, and possessing a strong, characteristic odour of Cocoa-nut

This substance is more suitable for the manufacture of (especially in the cooler months of the year) than Oil of Theobroma the latter is so near the temperature of the body that the mixtures made with it frequently take a very long time to melt Mixtures of Stearin and Theobroma Oil give intermediate figures

Tests—Cocoa-nut Stearin has a sp gr at 60° C (140° F) of about 0.896 It has a m p of 28.9° C (84° F) It possesses a Saponification value of about 256, and an Iodine absorption of about 5 When distilled by the Reichart-Vollney test the number of c c of Tenth-normal Volumetric Barium Hydroxide Solution required to neutralise the distillate should amount to about 3 c c

Not Official. STILLINGIA.

QUEEN'S ROOT

The Root of *Stillingia sylvatica*, L., is official in U.S. It is stated to contain an alkaloid 'Stillingine' which should not be confounded with the eclectic remedy 'Stillingin' Has been found useful in secondary syphilis, tuberculosis and cutaneous diseases

Fluid Extract (1 in 1), average dose 2 c c (30 minims), is official in U.S., and forms a convenient means of exhibition

STRAMONII FOLIA. STRAMONIUM LEAVES

Fr., STRAMOINE; GER., STECHAPFELBLÄTTER, ITAL., STRAMONIO, SPAN., ESTRAMONIO

The dried Leaves of *Datura Stramonium*, L.

They contain an alkaloid, Daturine, identical with Hyoscyamine

The Stramonium Leaves official in the U.S.P. are required to yield not less than 0.25 p c of mydriatic alkaloids

- **Medicinal Properties**—It is much used for spasmodic asthma in the form of **cigarettes** and **smoking mixtures**

Dose—*Ph Ger* maximum single dose, 0.2 gramme, maximum daily dose, 0.6 gramme

Under the title **Daturæ Folia**, the dried leaves of *Datura fastuosa*, L., var. *alba*, Nees, are official in the *Ind* and *Col Add*, for India, the Eastern and West Indian Colonies

Official Preparation—Tinctura Stramonii

Not Official—Extractum Stramonii, Fluidextractum Stramonii, Pulvis Stramonii Compositus, Folia Stramonii Nitrita, and Unguentum Stramonii

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr, Ger, Ital, Jap, Mex, Norw, Russ, Spai, Swed, Swiss and U S

Descriptive Notes—The Stramonium Leaves of commerce have, during the last few years, been frequently adulterated. The genuine leaves average about 3 to 6 in (7 to 15 cm) long, and 2 to 3 in (5 to 8 cm) broad, but are sometimes much larger, 9 by 6 in (22 by 15 cm), have an ovate outline, a petiole 1 in (25 mm) or more long, a sinuate margin with acute, rather distant triangular teeth of variable size, an unequal base, a minutely wrinkled surface when dried, a paler under surface, and a characteristic odour. The leaf has a bitterish, saline taste.

Under the microscope the powder is easily distinguished from that of Belladonna Leaves by the presence of cluster crystals and the absence of crystal-sand cells, by the epidermal cells not being striated, by the long hairs having rough or papillose walls and no terminal glands, and by the long palisade cells 5 or 6 times longer than broad, *Vogl Atlas*. None of the adulterants hitherto used possess these characters.

The leaves of *Datura Tatula*, L., have usually a purplish tinge on the petiole often extending to the midrib.

The leaves of *D. fastuosa*, L., var. *alba*, Nees, are official in the *Ind* and *Col Add*. The leaves are not acuminate, and are obtusely sinuate, but not dentate.

Tests—Stramonium Leaves yield from 10 to 15 p.c. of ash. It has been recommended that an ash limit of 15 p.c. and the microscopical characters should be given. The *USP* gives a method for the assay of the leaves which is identical with that for Belladonna Leaves given under *Belladonnæ Folia*. A weighed quantity of 10 grammes of Stramonium in No. 60 powder is employed for the determination.

Preparation

TINCTURA STRAMONII TINCTURE OF STRAMONIUM

1 of Stramonium Leaves, percolated with Alcohol (45 p.c.) to yield 5

BP '85 tincture was from the seeds, with Proof Spirit

Tincture of Stramonium *USP* is required to contain 0.025 p.c. w/v of mydriatic alkaloids from Stramonium. The *BP* Tincture is not standardised, the *Ph.G.* does not include a tincture.

Dose—5 to 15 minims = 0.3 to 0.9 c c

Official in Swiss, 1 of seeds in 10, *U.S.*, 1 of leaves in 10

Tests—Tincture of Stramonium has a sp gr of 0.950 to 0.964, it contains between 3 and 4 p c w/v of total solids and about 43 p c w/v of Absolute Alcohol. The *U.S.P.* assays the tincture by a similar process to that given for the assay of Fluid Extract of Belladonna Root given under Extractum Belladonnae Fluidum. 100 c c of the tincture are evaporated to about one-tenth its volume, sufficient Alcohol (94.9 p c) is added, if necessary, to dissolve any separated substance, and the assay then continued as directed. The final multiplication by 10 being omitted in the calculation, as 100 c c of the tincture and not 10 c c of the fluid extract are used for the operation.

Not Official

EXTRACTUM STRAMONII (*U.S.P.*)—The *U.S.P.* Extract is prepared by the evaporation, to a pilular consistence, of fluid extract of Stramonium at a temperature not exceeding 50° C (122° F). The extract is required to contain 1.0 p c of mydriatic alkaloids, and in the event of a greater alkaloidal content than the above being yielded when assayed, sufficient powdered Milk Sugar is added to reduce it to the standard.

Tests—The *U.S.P.* employs a method identical with that for the assay of Extract of Belladonna Leaves given under the heading of Extractum Belladonnae Viride. A weighed quantity of 5 grammes of the extract is employed for the determination.

FLUIDEXTRACTUM STRAMONII (*U.S.P.*)—An approximately 1 in 1 fluid extract prepared by exhausting Stramonium Leaves in No. 40 powder with a mixture composed of 2 volumes of Alcohol (94.9 p c) and 1 volume of Water, reserving the first percolate, evaporating the remainder at a temperature not exceeding 50° C (122° F) to a soft extract and dissolving this in the reserved percolate, finally adjusting the fluid extract to contain 0.25 p c of mydriatic alkaloids.

Tests—The Fluid Extract of Stramonium is assayed by the *U.S.P.* by a process identical with that employed for the *U.S.P.* assay of Fluid Extract of Belladonna Root given under Extractum Belladonnae Fluidum. A measured quantity of 10 c c of the Fluid Extract is employed for the determination.

PULVIS STRAMONII COMPOSITUS—Stramonium Leaves, Datuna Tatula, Cannabis Indica, and Lobelia Inflata, all in powder, of each 6 drms, Nitre, in powder, 1 oz, Eucalyptus Oil, 30 minims, mix thoroughly.

It burns well, gives off dense fumes, and affords great relief during asthmatic attacks.—*B.M.J.* '84, ii 465, '87, ii 494.

Several formulas, somewhat similar to the above, appear in the Hospital Pharmacopœias. Himrod's cure and several other similar preparations have also been recommended for asthma.

Folia Stramonii Nitrata—Coarsely powdered Stramonium Leaves, 2, Potassium Nitrate, 1, Water, 3, soak, and after 12 hours, dry.

UNGUENTUM STRAMONII—Extract of Stramonium, 10, Diluted Alcohol, 5, Hydrous Wool Fat, 20, Benzoinated Lard, 65,—*U.S.P.*
This has been incorporated in the *B.P.C.*

STRAMONII SEMINA

STRAMONIUM SEEDS

The dried ripe, black, reniform Seeds of *Datura Stramonium*, L

The mixed alkaloids of Stramonium are generally called **Daturine**, but are the same as contained in Belladonna, viz, a mixture of Hyoscyamine and Atropine

The dried pale brown, obovate, compressed Seeds of *Datura fastuosa*, L, var *alba*, Nees, are official in the *Ind* and *Col Add* for India and the Eastern Colonies, also **Tinctura Daturæ Seminum** (1 in 4), dose 5 to 15 minims = 0.3 to 0.9 cc

Medicinal Properties—Similar to those of Belladonna. Antispasmodic and sedative in spasmodic and bronchitic asthma. The Extract and the Tincture are used in convulsive cough as anti-spasmodics. The Extract has been given with success in hay asthma. Like Belladonna, it causes dilatation of the pupil.

Official Preparation—Extractum Stramonii

Not Official—Guttæ Daturinæ

Antidotes—Same as for poisoning with Belladonna, p. 224, also Morphine subcutaneously, and Chloroform Inhalation

Foreign Pharmacopœias—Official in Port (*Estramonio*), and Swiss

Tests—Stramonium Seeds yield from 2 to 3 p.c. of ash. The total alkaloids vary from 0.17 to 0.5 p.c., an average of 15 samples gave 0.35 p.c., in the Leaves the percentage of alkaloid varied from 0.32 to 0.47 p.c., an average of 11 samples being 0.38 p.c. The *USP* requires that the dried leaves contain not less than 0.25 p.c. of mydriatic alkaloids.

Preparation

EXTRACTUM STRAMONII EXTRACT OF STRAMONIUM

A firm Extract, prepared by exhausting Stramonium Seeds, in No. 40 powder, with Alcohol (70 p.c.), and evaporation of the percolate.

BP '85 used Proof Spirit and removed the fixed Oil by Ether.

Dose— $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Not Official

GUTTÆ DATURINÆ—Daturine Sulphate, 2 grains, Water, 1 fl. oz.—*London Ophthalmic*

Not Official

STRONTII BROMIDUM

STRONTIUM BROMIDE

$\text{SrBr } 6\text{H } 2\text{O}$, eq. 352.92

FR., BROMURE DE STRONTIUM, GER., STRONTIUMBROMID, ITAL., BROMURO DI STRONZIO, SPAN., BROMURO ESTRONCICO

Colourless, translucent, hexagonal, deliquescent prisms, having a bitter saline taste.

It should be kept in well stoppered glass bottles of a dark amber tint in a cool place and protected as far as possible from exposure to the air, as it has a tendency to deliquesce. *USP* states that it is also occasionally efflorescent.

Strontii Bromidum Exsiccatum is also commercial 69 of the anhydrous is equivalent to 100 of the crystallised salt

Solubility—2 in 1 of Water, 1 in 3 of Alcohol (90 p c) .

Medicinal Properties—Recommended in chronic gastritis and dilated stomach, in doses of 30 grains thrice daily, also in similar doses in epilepsy, is said by some to be less depressant than Potassium Bromide—*B M J* '92, ii 1286, '95, i 1089, 1252, *B M J E*, '95, i 76, *L* '92, i 47, '93, ii 46, '95, i 567, '96, ii 871, '98, ii 988 Many cases of epilepsy in which, if it gets a fair trial, it will have pre-eminence over other Bromide salts—*L* '07 i 20 In acute gastric catarrh, *P* ii 130, in vomiting, *T G* '93, 115, in enteritis, *M A* '95, 239, in exophthalmic goitre in children—*B M J* '98, ii 1042 Increases coagulability of blood—*L* '08, i 97

15 grains 3 times daily, increasing the amount if necessary to 20 grains and then to 30 grains and 40 grains 3 times daily until seizures are under control Whilst in some cases apparently of greater value than Potassium Bromide in controlling epileptic seizures, yet on account of the more rapid action of the latter, its more lasting effect, the smaller dose required, and lastly, its cheapness, the Potassium salt must be regarded as the more generally useful in the treatment of epilepsy—*L* '99, ii 411

It has an unpleasant, metallic taste

Dose—5 to 30 grains = 0.32 to 2 grammes

3 drms daily has been given for weeks without any unpleasant symptoms—*L* '98, ii 988

Foreign Pharmacopœias—Official in Fr, Mex, Span and U S

Tests—Strontium Bromide melts when heated, and finally loses all its Water of crystallisation, equivalent to 30.4 p c The *U S P* anhydrous salt fuses at 690° C (1266° F) A crystal of the salt Hydrochloric Acid and introduced in a loop of platinum wire into flame gives a brilliant crimson coloration The salt dissolves readily in Water, forming a clear solution, which should be neutral in reaction towards Litmus A 5 p c aqueous solution affords on the addition of a saturated solution of Calcium Sulphate a white precipitate insoluble in diluted acids Potassium Chromate Solution affords a yellow precipitate soluble in Acetic Acid, Ammonium Carbonate Solution yields a white precipitate soluble with effervescence in Acetic Acid The aqueous solution affords with Silver Nitrate Solution a yellowish curdy precipitate practically insoluble in Ammonia Solution The percentage of pure Strontium Bromide present in a specimen may be determined by titrating a weighed quantity of the salt with Tenth-normal Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator of neutrality The *U S P* requires that a weighed quantity of 0.5 of a gramme when dissolved in about 50 c c of Water shall require not less than 27.4 (27.48) c c nor more than 29.4 c c of the volumetric solution to produce a permanent red colour, corresponding to at least 97 p c of pure Strontium Bromide

The more generally occurring impurities are excess of Water, Copper, Lead, Barium, Chlorides and Iodides The aqueous solution either when rendered faintly alkaline with diluted Hydrochloric Acid, or when rendered faintly alkaline with Ammonia Solution, should not be darkened in colour by Hydrogen Sulphide indicating the absence of Copper, Lead and Zinc A weighed quantity of 1 gramme of the salt when mixed with an equal weight of Sodium Acetate, and the mixture dissolved in 5 c c of Distilled Water when rendered faintly acid with 3 to 5 drops of diluted Acetic Acid and mixed with 5 drops of Potassium Chloride Solution should not afford a cloudiness within 3 minutes, indicating the presence of Barium If the precipitate obtained by completely precipitating an aqueous solution of the salt with Silver Nitrate Solution be separated, shaken with Ammonia Solution and filtered, upon adding Nitric Acid in slight excess, to the filtrate, no pronounced turbidity should result, indicating the absence of more than traces of Chloride If a little Chlorine Water diluted with an equal volume of Water be added carefully drop by drop to 10 c c of an aqueous

5 p.c. solution of the salt and the aqueous mixture be shaken with a little Carbon Bisulphide, the Carbon Bisulphide solution should assume a yellow or yellowish brown colour, and should be free from any violet tint, indicating the absence of Iodides

STRONTII CINNAMAS —A white, or whitish, amorphous powder, soluble 1 in 100 of Water, insoluble in Alcohol (90 p.c.) It has been used suspended in 3 parts of Glycerin to 5 parts of Water in malignant disease —*L.* '03, 11 750

Tests —Strontium Cinnamate when strongly heated evolves an aromatic odour resembling Benzaldehyde, and finally burns leaving a carbonaceous residue which when dissolved in Water possesses a strongly alkaline reaction towards red Litmus paper, and effervesces on the addition of diluted Hydrochloric Acid, the solution when neutralised answering the tests distinctive of Strontium given under the heading of Strontii Bromidum. The Cinnamic Acid separated from the salt should possess the m.p. and answer the tests distinctive of Cinnamic Acid given under the heading of Acidum Cinnamicum

STRONTII IODIDUM ($\text{SrI}_2 \cdot 6\text{H}_2\text{O}$, eq. 446.02) —Translucent, colourless, hexagonal prisms or a white granular powder, readily soluble in Water, possessing a bitter saline taste. It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with air, as it is of a deliquescent nature and liable to change to a yellow colour on exposure to the light. It has been used in place of the alkali Iodides in chronic endocarditis

Dose — $\frac{7}{8}$ to 15 grains = 0.5 to 1 gramme

Foreign Pharmacopœias —Official in U.S.

Tests —The salt melts when cautiously heated, gradually losing the whole of its Water of crystallisation, equivalent to 24.05 p.c., and leaving a residue of the anhydrous salt. It dissolves in Water, forming a clear solution which is neutral to Litmus, or but very faintly alkaline to red Litmus paper. It yields the tests distinctive of Strontium given under the heading of Strontium Bromide. The aqueous solution affords with Silver Nitrate Solution a yellow curdy precipitate insoluble in Nitric Acid, insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution. The aqueous solution when cautiously mixed with diluted Chlorine Water affords a yellowish coloration changing to blue on the addition of Mucilage of Starch. If the liquid before the addition of Starch solution be shaken with Carbon Bisulphide, the Carbon Bisulphide Solution assumes a strong violet colour. The percentage of pure Iodide present in a specimen may be determined by titrating a weighed quantity with Tenth normal Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator. The *U.S.P.* employs an excess of Tenth normal Volumetric Silver Nitrate Solution, titrating the excess with Tenth-normal Volumetric Potassium Sulphocyanate Solution, using Ferric Ammonium Sulphate T.S. as an indicator. A weighed quantity of 0.5 of a gramme of the Iodide is dissolved in about 100 c.c. of Distilled Water, 25 c.c. of Tenth normal Volumetric Silver Nitrate Solution added, 5 c.c. of Nitric Acid and 5 c.c. of Ferric Ammonium Sulphate Solution, the flask stoppered and shaken, not less than 17 c.c. nor more than 31 c.c. of Tenth normal Volumetric Potassium Sulphocyanate Solution should be required to produce a permanent red tint, corresponding to 98 p.c. of pure Strontium Iodide.

The more generally occurring impurities are Copper, Lead, Iron and Zinc, Barium and Chlorides. Neither the aqueous solution slightly acidified with diluted Hydrochloric Acid nor an aqueous solution made faintly alkaline with Ammonia should afford any darkening in colour on the addition of Hydrogen Sulphide, indicating the absence of Copper, Lead, Iron and Zinc. The presence of Barium may be detected by a test similar to that described under Strontium Bromide. If the precipitate obtained by completely precipitating an aqueous solution of the salt with Silver Nitrate, be separated, shaken with Ammonia and filtered, the filtrate rendered faintly acid with Nitric Acid should not yield any pronounced turbidity, indicating the absence of more than traces of Chlorides.

STRONTII LACTAS ($\text{Sr}(\text{C}_2\text{H}_3\text{O}_2)_2 \cdot 3\text{H}_2\text{O}$, eq. 317.32) —A white granular powder, or in crystalline nodules, soluble 1 in 3 of Water. Has been recommended

for albuminuria in parenchymatous nephritis —*L* '92, i 47, 3, i 567, 96, i 255, *T G* '94, 461, *B M J P* '96, i 76, '97, ii 40. Excellent diuretic in Bright's disease —*L* '94, ii 992

Dose —20 to 30 grains = 1.3 to 2 grammes

Official in Fr and Mex

Tests —Strontium Lactate when heated to 110° C (230° F) loses its Water of crystallisation, equivalent to 16.9 p.c. At a still higher temperature it yields inflammable vapours, and burns leaving a carbonaceous residue which effervesces strongly on the addition of Hydrochloric Acid and which produces a distinctive crimson flame test of Strontium. The salt dissolves in Water, yielding a clear solution which is slightly acid in reaction towards blue Litmus paper. It answers the tests distinctive of Strontium given under the heading of Strontium Bromidum. The aqueous solution when acidified with Sulphuric Acid and treated with Tenth-normal Volumetric Potassium Permanganate Solution decolorises the Permanganate, evolving simultaneously an odour of Aldehyde. The percentage of pure Strontium Lactate may be determined by extracting the carbonaceous residue left on igniting the carefully dried salt, with boiling Water, until the washings no longer effect Methyl Orange Solution, the filtrate and washings being titrated with Normal Volumetric Sulphuric Acid Solution, using Methyl Orange Solution as an indicator of neutrality. A weighed quantity of 1.33 grammes of the salt thus treated should require for complete neutralisation not less than 9.9 c.c. of the Normal Volumetric Acid Solution, corresponding to at least 98.6 p.c. of the pure salt.

The more generally occurring impurities are Arsenic, Copper, Lead, Iron and Zinc, Barium, Carbonates and Oxalates, Chloride, Butyrate, Propionate, and readily charred organic impurities. Arsenic, Copper, Lead, Iron and Zinc may be tested for as described under Strontium Bromide or Strontium Iodide, as also may Barium. The 5 p.c. aqueous solution of the salt should be perfectly clear leaving no weighable residue, and no effervescence should occur on mixing 0.5 of a gramme of the salt with 1 c.c. of Sulphuric Acid, indicating the absence of Carbonates and Oxalates. The aqueous solution of the salt should not afford more than 0.5 c.c. of a precipitate with Silver Nitrate Solution, indicating the absence of traces of Chloride. A solution of the salt in concentrated Sulphuric Acid should be free from perceptible or penetrating odour, even at heating, indicating the absence of Butyrate and Propionate, and this Sulphuric Acid Solution should not become in 10 minutes more than a pale straw yellow colour, indicating the absence of readily charred organic impurities. The salt should not lose, when carefully dried at the temperature mentioned above, more than its proper quantity of Water of crystallisation.

STRONTIUM SALICYLAS ($\text{Sr}(\text{C}_7\text{H}_5\text{O}_2)_2 \cdot 2\text{H}_2\text{O}$, eq 394.72) —A white powder, slightly soluble in Water.

It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light.

Has been recommended as an intestinal antiseptic, also in gouty and rheumatic conditions —*C D* '95, i 291, *P J* '96, ii 63, '97, ii 118.

Dose —5 to 15 grams = 0.32 to 1 gramme

Official in U S

Tests —Strontium Salicylate decomposes on heating, giving off inflammable vapours and an odour of Phenol, and leaving a residue of Strontium Carbonate. This residue dissolves with effervescence in Hydrochloric Acid, giving a solution which affords the distinctive flame test for Strontium. The salt dissolves in Water, forming a clear solution which is faintly alkaline in reaction towards red Litmus paper. This aqueous solution answers the tests distinctive of Strontium given under Strontium Bromidum. A 1 p.c. aqueous solution of the salt affords with Ferric Chloride Test a deep violet coloration. A 5 p.c. aqueous solution affords with Copper Sulphate Test a green coloration. A small quantity of the salt when warmed with a few drops of concentrated Sulphuric Acid and a little Methyl Alcohol evolves the distinctive odour of Methyl Salicylate.

The *U S P* method of determination is a gravimetric one, 0.5 of a gramme of the salt is moistened with 1 c.c. of Sulphuric Acid, the mixture cautiously heated until no more vapours are given off, the residue again moistened with a few drops of the acid, again heated and finally ignited until of a constant weight. The *U S P* requires that the residue of Strontium Salicylate should weigh not less than 0.227 gramme corresponding to not less than 98.5 p.c. of pure Strontium Salicylate. The Salicylic Acid separated by acidification from the aqueous solution, washing the precipitate till free from mineral acid and carefully drying, should possess the m.p. and answer the tests distinctive of Salicylic Acid given under Acidum Salicylicum.

The more generally occurring impurities are Copper, Lead, Iron and Zinc, Barium, Carbonates and Chlorides. The same methods may be employed for the detection of Copper, Lead, Iron, Zinc and Barium as are employed in the examination of Strontium Bromide and Strontium Iodide. The concentrated aqueous solution should afford a white crystalline precipitate but yield no effervescence on the addition of diluted Nitric Acid, indicating the absence of Carbonate, and if the crystalline precipitate be filtered off, the filtrate should not afford a distinct turbidity on the addition of Silver Nitrate Solution, indicating the absence of more than traces of Chlorides. The absence of excess of Water of crystallisation may be ensured by a determination of the loss of weight on heating the salt at 100° C (212° F).

STROPHANTHI SEMINA

STROPHANTHUS SEEDS

FR, STROPHANTHUS, GLR, STROPHANTHUSSEMIN, ILL, STROPHANTHUS
SPAN, ESTROPHANTHUS

The dried ripe Seeds of *Strophanthus Kombé*, freed from the awns

The commercial seed usually contains the seeds of other species in addition to those of *S. Kombé*—*P J* (3) 219 660. The active principle is a glucoside, *Strophanthin*.

Medicinal Properties—A cardiac tonic. Especially valuable in mitral regurgitation with failure of compensation, and in aortic regurgitation accompanied by cardiac insufficiency. The active principle being very soluble and diffusible, *Strophanthus* acts with such rapidity that it is more useful than *Digitalis* in promptly stimulating extreme or sudden cases of cardiac failure. Of great value in avoiding both the cardiac embarrassment so frequently fatal in acute pneumonia and the collapse which may occur at the crisis. It is easily eliminated, it is not cumulative, it can be administered over a long period of time, and, unless there be marked gastrointestinal catarrh, it has no tendency to produce digestive disturbance. It has acted beneficially in many cases in which *Digitalis* has failed or has disagreed.

Strophanthus acts more energetically on the heart than on the vessels, whereas *Digitalis* acts on the vessels as much as, or even more than, on the heart. *Digitalis* thus possesses the power of increasing arterial tension, and so of putting extra strain on the heart, therefore, in those cases in which pulse tension is high, *Strophanthus* is to be preferred.

A more powerful cardiac tonic than Digitalis and superior as a diuretic — *B M J* '95, i 368, *B M J E* '97, ii 3, '98, i 12, *T G* '98, 36

In Graves' disease — *L* '93, ii 822 In alcoholism — *L* '94, ii 212

As to the disparity in the results obtained by different observers, Fraser remarks that, 'there are several species of the genus, and that while the therapeutic effects have been determined with only one of these species, the seeds of several of the others have indiscriminately been substituted. The whole fruit, and not the seeds only, and immature seeds, poor in the active principle and rich in irritating resin, have been used to prepare the Tincture seeds already exhausted with Alcohol have been re-sold in the market, and further, even when good seeds were used, Petroleum Ether has been substituted for Ethylic Ether, preparatory to percolation with Rectified Spirit, with the result that the Tincture (1865) contained much resin, which produced stomach and intestinal disorder.'

Official Preparations — Extractum *Strophanthi* and Tinctura *Strophanthi*

Not Official — *Strophanthum* and *Ouabain*

Foreign	Official in Austri, Belg, Dan, Dutch, Fr, Ger, Ital, Spain, Swiss and U S
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Descriptive Notes.—For many years past the *Strophanthus* seeds of commerce have been almost invariably a mixture derived from different species. The official seeds are limited to those of *Strophanthus Kombé*, Oliver. They are about $\frac{5}{8}$ in (15 mm) long and rather more than $\frac{1}{2}$ in (4 mm) broad (15 mm long, 4 to 5 mm wide and 2 to 2.5 mm in thickness, *USP*, 17 mm long, 5 mm broad, 3 mm thick, *PG*), of a greenish-fawn colour and covered with silky adpressed hairs, linear, elliptical, acuminate, compressed, rounded at the base and having a longitudinal ridge on one side from $\frac{1}{2}$ in below the centre to the apex of the seed, the cotyledons are straight, surrounded with a thin endosperm, the odour is characteristic, the taste very bitter.

The seeds vary a little in size and shape according to their position in the pod, the lowest being usually rather longer, more acuminate, and furnished with longer awns.

The pods and seeds of *Strophanthus Courmontii*, Sacl, var *fallax*, Holmes, so closely resemble in size, shape and colour those of *S. Kombé* that it is practically impossible to distinguish them by sight, although the flowers and leaves of the two species are quite distinct. There are microscopical differences between the two seeds, but it needs an expert to detect them.

The colour of the seeds appears brownish or according to the incidence of the light and the observer with regard to it, due to the disposition of the hairs. But when a section of the seed is immersed in a mixture of 8 parts concentrated Sulphuric Acid (*BP*) strength and 2 of Water, a deep green colour rapidly appears in the albumen surrounding the embryo to which the colour gradually extends. The seeds of no other known species except those of *S. hispidus*, DC give this reaction, and the seeds of that species are smaller and dark brown in colour. They also contain the same active principle, but just as there are seeds closely resembling those of *S. Kombé*, which slowly give a pink reaction, so there is a seed closely resembling *S. hispidus*, viz, *S. Arnoldianus*, De Wild and Dur, in its brown colour and size which gives a pink reaction, so that in either case a colour test is necessary. Several of

the species are not regarded by the African natives as poisonous, but the woolly Zambesi seeds, *S. Nicholsoni*, Holmes, that have appeared in commerce, distinguished by the longer white hairs hiding the apex of the seed, and those of *S. gratus*, Fianch, are recognised by the natives as poisonous and are used in making arrow poison. The seeds of the latter are the only W African kind imported into commerce that are quite hairless. Both of these give a pink coloration with Sulphuric Acid, but it is probable that this reaction, though useful to distinguish other seeds from those of *S. Kombé*, does not always depend upon the presence of Ouabain as it does in the seeds of *S. gratus*. At the present time it is possible to obtain commercially in London the seeds of *S. Kombé* unmixed with other kinds. It should be noted that the acid used for testing is apt to become weaker by keeping, and recently mixed acid should therefore be employed, as a weak acid does not readily give the green reaction. The *PG* points out that the starch grains in the official seed do not exceed 0.008 mm, and the *USP* that the hairs appear under the microscope of a light brownish green colour, are thin walled, 1-celled, and 1 mm or less in length.

Tests—*Strophanthus* seeds contain from 3 to 4 p.c. of ash, and 5 p.c. is rarely exceeded. The *BP* states that Sulphuric Acid colours the endosperm and sometimes the cotyledons a dark green, indicating the presence of *Strophanthin*. The *BP* employs a concentrated acid. The author found from an examination of commercial tinctures of *Strophanthus* that the use of a Sulphuric Acid slightly diluted with Water gave a more definite reaction. Holmes has suggested (*PJ* '02, i 254) that in a future edition of the *BP* the words 'Sulphuric Acid' should be replaced by 'a freshly prepared solution containing 8 parts of Sulphuric Acid and 2 of Distilled Water'. Several processes have been suggested for the chemical assay of *Strophanthus*, but not one of them is entirely satisfactory. The following process described (*PJ* '96, ii 463, '02, ii 281, 304) is easy of manipulation and yields results which possess a certain value as a criterion of the activity of the preparation. It is based upon the determination of the amount of *Strophanthidin* produced on the hydrolysis of *Strophanthin*. A measured quantity of 50 c.c. of the tincture is diluted with 50 c.c. of Water and the Alcohol removed by distillation. The filtered aqueous liquid, after being shaken with Chloroform, is digested for 1 hour on a water-bath with diluted Sulphuric Acid, after cooling, the turbid liquid is agitated with 3 successive small quantities of Chloroform, the chloroformic layer in each case is separated, transferred to a tared flask, the Chloroform removed by evaporation, the residue of *Strophanthidin* dried below 65.6° C (150° F) and when constant weighed. The percentage of *Strophanthidin* found divided by 0.365 corresponds to the percentage of *Strophanthin* present. The physiological method of assay is also in vogue, but it is doubtful whether, in the present state of knowledge, the physiological process possesses any advantages over the chemical. Tinctures made from

the seeds answering the official requirements, when carefully prepared, have yet to be shown wanting in activity. So-called physiological standardisation of galenicals leaves much to be desired. A method for rapidly and approximately estimating Strophanthin in the extract and tincture by the optical rotation has been suggested (*C D* '98, ii 289)

Preparations

EXTRACTUM STROPHANTHI EXTRACT OF STROPHANTHUS

1 of Strophanthus Seeds, exhausted with Purified Ether, and dried, then percolated with Alcohol (90 p c) until 10 of percolate is obtained, concentrate this by evaporation to a thick liquid, and add Milk Sugar *q s* to yield 2 of Extract, in powder.

Dose.— $\frac{1}{4}$ to 1 grain = 0.016 to 0.06 gramme

Official in Mex

TINCTURA STROPHANTHI TINCTURE STROPHANTHUS

Percolate 1 of Strophanthus Seeds, in No. 30 powder, with Alcohol (70 p c) until 20 is obtained, and dilute with Alcohol (70 p c) to yield 40 (1 in 40)

B P 1898 reduced the strength from 1 in 20 to 1 in 40, making the dose uniform with Tincture of Digitalis

Dose.—5 to 15 minims = 0.3 to 0.9 c c

Foreign Pharmacopœias—Official in Ital, 1 in 20, Norw, 1 and 10, Austr, Belg, Dan, Dutch, Fr, Ger, Jap, Russ, Span, Swed, Swiss and U S, 1 in 10, Mex, 1 in 5. All by weight except U S. The *Brussels Conference* agreed to a strength of 10 p c, prepared by percolation with Alcohol (70 p c), the seeds not to be freed from fat.

Tests.—Tincture of Strophanthus has a sp gr of 0.890 to 0.894, it contains from 0.4 to 0.9 p c w/v of total solids and about 68 p c w/v of Absolute Alcohol. 2 c c of the tincture evaporated on a water-bath and the residue, when cool, moistened with a drop of a freshly prepared solution (0.1 in 100) of 8 parts of Sulphuric Acid and 2 of Distilled Water should yield a green and not a red coloration. A standard of 0.2 p c has been suggested for the tincture. A method of determining the Strophanthin is described under Strophanthus Semina. Eight samples of tincture obtained from houses of the highest repute examined in the author's laboratory showed percentages of Strophanthin varying from 0.082 to 0.301 p c, with an average of 0.109 p c. When tested by the Sulphuric Acid test, as described above, only one sample (that of Fiasel's) gave a green coloration, the others yielding either a yellow or purple coloration. Two samples of the *B P* tincture prepared in the author's laboratory both gave pure green colorations.

Not Official.

STROPHANTHIN—A pale yellow amorphous powder, or in white microscopic crystalline plates. It possesses an agreeable, bitter taste and is extremely poisonous.

Solubility—Freely in Water and in Alcohol (90 p c), practically insoluble in Chloroform, Ether, and in Carbon Disulphide.

Dose — $\frac{1}{400}$ to $\frac{1}{200}$ grain — 0.0002 to 0.00032 grammes

It is official in the *U S P* and is described as a glucoside or mixture of glucosides obtained from *Strophanthus*. It is also official in the *F. Codex* and *Mex*.

It should be kept in well stoppered glass bottles of a dark amber tint and exposed as little as possible to the light.

Tests —*Strophanthin* when heated darkens at about 146° C (294.8° F), becomes pasty at 165° C (329° F), and melts at 172.5° C (342.5° F). The *U S P* states that it commences to fuse at 170° C (338° F), and is not completely melted until the temperature reaches 190° C (374° F). *F. Codex* (1908) gives the melting point as 185° C (365° F). It dissolves readily in Water, the solution being neutral in reaction to Litmus paper and dextrogyrate. A trace of the substance when moistened with a freshly prepared solution containing 8 parts of Sulphuric Acid and 2 of Distilled Water yields an emerald green coloration, subsequently changing to brown. An aqueous solution yields on the addition of a trace of Ferric Chloride T.S. and a few cc of Sulphuric Acid a reddish brown precipitate, turning dark green after 1 or 2 hours. Tannic Acid Solution throws down from an aqueous solution a copious white precipitate, which redissolves on agitation until an excess of the reagent has been added. The usual reagents employed as alkaloidal precipitants, e.g., Potassio mercuric Iodide Solution and Iodo-potassium Iodide Solution, produce no precipitate in solutions of the glucoside. Potassio cupric Tartiate (Fehling's) Solution produces no red precipitate of Cuprous Oxide when boiled with the aqueous solution, but if an aqueous solution be heated to 70° C (158° F) with a small amount of Diluted Sulphuric Acid, the *Strophanthin* undergoes hydrolysis with the formation of Glucose and *Strophanthidin*, the latter separating out as a flocculent precipitate which can be filtered off, the filtrate if boiled with Potassio cupric Tartiate (Fehling's) Solution will now yield a red precipitate of Cuprous Oxide. When ignited with free access of air it should leave no weighable residue.

Ouabain —White odourless and tasteless slender transparent needles, practically insoluble in cold Water and Alcohol (90 p.c.) insoluble in Ether and in Chloroform. Recommended medicinally in half the doses of *Strophanthin*, in cases where *Strophanthus* and *Digitalis* both fail. The action is similar to *Strophanthin*. It is obtained from *Acokanthera Schumperi*, Oliv. (*S. Ouabaro*, Cath.), of the same natural order as *Strophanthus*, and from the seeds of *S. gratus*.

STRYCHNINA.

STRYCHNINE

$C_{21}H_{22}N_2O_2$, eq 331.75

Translucent, colourless rhombic prisms or a white crystalline powder. Permanent in the air. This alkaloid is odourless, but possesses an intensely bitter taste and is extremely poisonous. It may be obtained from the dried ripe seeds of *Strychnos Nux Vomica*, *Ignatia Amara*, and other species of *Strychnos*.

Strychnine is official in the *B P* and *U S P*, but not in the *P G*.

Solubility —1 in 6000 to 8000 of Water, 1 in 170 of Alcohol (90 p.c.), 1 in 250 of Alcohol (70 p.c.), about 1 in 400 of Alcohol (60 p.c.), 1 in 800 of Alcohol (45 p.c.), 1 in 4200 of Alcohol (20 p.c.), 1 in 350 of Absolute Alcohol, 1 in 6 of Chloroform, nearly insoluble in Ether.

Medicinal Properties —Similar to those of *Nux Vomica*, gastric, cardiac, and general tonic, useful in the treatment of reflex

or functional paralysis, and of peripheral neuritis and paralysis due to Alcohol, tobacco, or diphtheria, also in cases of lead-palsy. It increases peristalsis, and is therefore a useful addition to other purgatives. Recommended in chronic alcoholism, muscular tremors, tobacco amblyopia, impotence and nervous exhaustion. For other uses and for its contra-indications, see *Nux Vomica*. It has a cumulative action and is a very active poison.

An antidote in Chloroform poisoning — *B M J E* '94, 1 47. In snake bites — *T G* '93, 542, '94, 517.

In the treatment of surgical shock 10 minims of Liquor Strychninæ given subcutaneously just before commencing anæsthesia, followed after the operation by 5 minims subcutaneously injected every 2 hours for several hours if called for — *L* '02, 1 1025, 1063, 1210, 1357, 1497, *B M J* '99, 11 1471.

In diabetes insipidus good results are stated to have followed the injection of $\frac{1}{10}$ grain of the Nitrate on each of the first 2 days, and $\frac{1}{2}$ grain on the third — *B M J E* '04, 11 71, *B M J E* '06, 11 72.

Of undoubted value in collapse from cardiac weakness following infantile diarrhoea. Best administered in doses of $\frac{1}{2}$ minim of the Liquor Strychninæ — *Pr* lxxv 508.

In beri-beri $\frac{1}{10}$ grain administered twice daily caused a rapid improvement in a week — *L* '05, 11 540.

In surgical shock, notwithstanding an article which appeared by a distinguished authority stating its administration in large doses hypodermically was stated to be of very little value (*L* '05, 1 780) in the treatment, and in many cases it directly contributed to a fatal issue.

May be made use of to prevent surgical shock if it can be administered in small doses at reasonable intervals for, say, a week or 10 days previous to an operation — *L* '05, 1 851.

In the treatment of shock, stimulants, and especially Strychnine, are absolutely contra-indicated, as they tend to increase the severity of the condition and to retard recovery. Adrenalin, Hemisine, or Ergot are recommended — *L* '05, 1 854.

To the exhausted anæmic, or overworked debilitated person, Strychnine, 5 to 7 minims of the liquor, is the hypnotic *par excellence* — *F T* '07, 70.

Morphomania treated successfully by Atropine and Strychnine — *B M J* '07, 1 1173.

Strychnine, especially in form of injection and *Nux Vomica*, are powerful heart tonics — *B M J* '06, 11 987.

Dose. — $\frac{1}{60}$ to $\frac{1}{15}$ grain = 0.0011 to 0.0044 gramme.

Prescribing Notes. — May be given in the form of pill well triturated with Milk Sugar and the addition of 'Diluted Glucose,' q s, but it is more frequently prescribed in solution.

Antidotes. — Animal Charcoal or Tannic Acid, followed by an emetic, or the stomach-tube. Potassium Bromide, in $\frac{1}{2}$ oz in Water, with 30 grains of Chloral 2 dim of the Bromide, with or without 10 grains of Chloral, may be given every 15 or 20 minutes if necessary. Amyl Nitrite inhalations, the Amyl being poured freely on a handkerchief and held close to the nose. The patient may be kept fully under Chloroform or Ether. Curare, $\frac{1}{2}$ grain, by hypodermic injection. Antidote reserved, if possible — *Murrell*.

A case of recovery after taking 3 grains of Strychnine — *L* '67, 11 41, 118.
S gran. of Morphine to be an antidote for 1 grain of Strychnine — *L* '71, 11, 540.

Foreign Pharmacopœias. — Official in Fr, Port, Mex, Span and U S.

Tests. Strychnine melts at 265° to 266° C (509° to 510° F), the U S P says 265° C (514° F). It dissolves very sparingly in Water, the aqueous solution being alkaline towards red Litmus paper and being lævogyrate. Even in highly dilute solution it possesses an

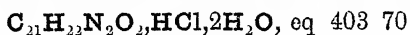
extremely bitter taste, but the solution should be tasted with extreme caution. A crystal moistened with Sulphuric Acid produces a colourless solution, which, on the addition of a minute crystal of Potassium Bichromate, assumes an intense purple-violet coloration, passing from red to yellow. A similar coloration is produced when Sulphuric Acid containing a one thousandth part of Potassium Permanganate is brought into contact with a crystal of the alkaloid, but the rotation of tints is very rapid and the reagent itself is apt to give a more or less purple-violet colour with Sulphuric Acid. With a drop of Sulphuric Acid containing a trace of Ammonium Vanadate (1 gramme of Ammonium Vanadate in 100 cc) the alkaloid produces a deep purple violet coloration. Sulphuric Acid containing a trace of Potassium Iodate also produces with the alkaloid a purple-violet coloration, changing to reddish purple. The free alkaloid may be determined by titration with Tenth normal Volumetric Sulphuric Acid Solution, using Iodeosin Solution as an indicator of neutrality, 1 cc of Tenth-normal Volumetric Sulphuric Acid is equivalent to 0.033175 gramme of Strychnine.

The more generally occurring impurities are Brucine, Sugar and other readily charred organic impurities, and mineral impurities. The alkaloid should not be coloured on the addition of concentrated Nitric Acid, indicating the absence of Brucine. It should dissolve in cold concentrated Sulphuric Acid without alteration in colour, indicating the absence of Sugar and readily charred organic impurities. When ignited with free access of air it should leave no weighable residue.

STRYCHNINÆ HYDROCHLORIDUM.

STRYCHNINE HYDROCHLORIDE

HYDROCHLORATE OF STRYCHNINE — *B.P.* '85



Translucent, colourless, prismatic crystals or white silky crystalline needles, it is efflorescent in dry air, and should therefore be kept in well-stoppered bottles. It possesses an intensely bitter taste. It is officially described as the Hydrochloride of an alkaloid obtained from *Nux Vomica* and from other species of *Strychnos*, but would have been preferably described as the Hydrochloride of the alkaloid Strychnine.

Strychnine Hydrochloride is only official in the *B.P.*, in the *U.S.P.* both the Nitrate and the Sulphate are official, in the *P.G.* only the Nitrate is official.

Solubility — 1 in 35 Water, 1 in 73 Alcohol (90 p.c.), insoluble in Ether.

Medicinal Properties — See 'Strychnina'.

Dose — $\frac{1}{40}$ to $\frac{1}{15}$ grain = 0.0011 to 0.0044 gramme.

Ph. Ger. maximum single dose, 0.01 gramme, maximum daily dose, 0.02 gramme of the Nitrate.

Prescribing Notes—*In solution, tablet or pill* A good pill is made by well triturating with Milk Sugar and massing with 'Diluted Glucose' Strychnine is usually given immediately after a meal Solution of Strychnine is frequently prescribed with Solution of Arsenic, in which case the Liquor Arsenici Hydrochloricus should be ordered and not the Alkaline Liquor

Official Preparation—Liquor Strychninæ Hydrochloridi

Not Official—Mistura Strychninæ Acida, Strychninæ Nitras, Strychninæ Sulphas, Strychnine Acetate, Strychnine Hydrobromide, Strychnine Valerianate

Incompatibles—Alkalis and Alkaline Carbonates, Bromides and Iodides, Liquor Sodii Arsenatis, and Liquor Arsenicalis

Foreign Pharmacopœias—Official in Mex

Tests—Strychnine Hydrochloride when heated loses its Water of crystallisation, slowly and incompletely at 100° to 110° C (212° to 230° F), readily and completely at 130° to 135° C (266° to 275° F) If of the pharmacopœial formula, it should lose theoretically 8.8 p.c., but the balance of opinion seems to be that a salt containing the full amount of Water of crystallisation is not a commercial article The Report of the Committee of Reference in Pharmacy does not go far in elucidating the matter, it is content with stating that the composition of the salt needs re-investigating, and that the temperature at which this salt is stated to lose its Water of crystallisation is too low The B.P. authorities themselves do not appear to be over-confident of the correctness of the formula, as they state under the heading of Tests that when dried at a temperature of 100° C (212° F) it should lose from 7.3 to 8.8 p.c. of moisture Commercial specimens of the salt lose about 7.3 p.c. of Water at temperatures between 100° and 130° C (212° and 266° F) The salt dissolves in Water, forming a clear solution, which should be neutral in reaction towards Litmus paper It should afford on the addition of Ammonia Solution a white precipitate soluble in Ether If the ethereal solution be separated, evaporated to dryness, it should yield a residue which answers the tests distinctive of Strychnine given under that heading An aqueous solution of the salt acidified with Nitric Acid yields on the addition of Silver Nitrate Solution a white, curdy precipitate, which when separated, washed and treated with Ammonia Solution, dissolves and is again reprecipitated on acidification with Nitric Acid An aqueous solution of the salt when acidified with diluted Hydrochloric Acid should yield no turbidity on the addition of Barium Chloride Solution, indicating the absence of Sulphates

Preparation

LIQUOR STRYCHNINÆ HYDROCHLORIDI SOLUTION OF STRYCHNINE HYDROCHLORIDE SOLUTION OF HYDROCHLORATE OF STRYCHNINE—B.P. '85

Strychnine Hydrochloride, $17\frac{1}{2}$ grains, Alcohol (90 p.c.), 1 fl. oz., Distilled Water, $q.s.$ to yield 4 fl. oz. (1 in 100)

Dose—2 to 8 minims = 0.12 to 0.5 c.c.

11 minims contain $\frac{1}{10}$ grain of Strychnine Hydrochloride
2 minims subcutaneously " " " " " " " "

Not Official

MISTURA STRYCHNINÆ ACIDA—Solution of Strychnine Hydrochloride, 3 minims, Diluted Nitro Hydrochloric Acid, 15 minims, Glycerin, 30 minims, Compound Infusion of Gentian, to 1 fl oz—*St Thomas's*

This has been incorporated in the *B P C*

STRYCHNINÆ NITRAS Strychnine Nitrate ($C_{15}H_{21}NO \cdot HNO_3$, eq 394.93)—Colourless, silky, crystalline needles, possessing an extremely bitter taste. It is the Nitrate of the alkaloid Strychnine, and is official in the *U S P* and *P G*, but not in the *B P*

It should be kept in well stoppered glass bottles

Solubility—1 in 63 of Water and 1 in 120 of Alcohol (90 p c)

Official in Aust, Belg, Dan, Dutch, Ger, Hung, Ital, Jap, Mex, Norw, Russ, Swed, Swiss and U S

Tests—Strychnine Nitrate when heated decomposes. It dissolves in Water, forming a clear solution possessing even when highly diluted an extremely bitter taste (it should be used with extreme caution), which is neutral in reaction towards Litmus paper and Levogyrate. When heated with Hydrochloric Acid a bright red coloration is produced. The alkaloid separated from an aqueous solution of the salt by precipitation, and solution by an immiscible solvent should answer the tests distinctive of Strychnine given under that heading. A solution of the salt poured carefully upon Sulphuric Acid containing a little Diphenylamine develops a blue coloration at the junction of the two liquids. The salt when moistened with Sulphuric Acid should not assume more than a faint yellow colour, indicating the limit of Brucine. It should leave no weighable residue when ignited with free access of air, indicating the absence of mineral impurities.

Hypodermic Tablets are made containing $\frac{1}{100}$ and $\frac{1}{1000}$ grain Strychnine Nitrate, and $\frac{1}{30}$, $\frac{1}{60}$, $\frac{1}{100}$, and $\frac{1}{150}$ grain of Strychnine Sulphate

STRYCHNINÆ SULPHAS Strychnine Sulphate [$(C_{15}H_{21}NO) \cdot H_2SO_4 \cdot 5H_2O$, eq 850.24]—Colourless, or white, odourless, efflorescent, prismatic crystals, possessing an intensely bitter taste. It is the Sulphate of the alkaloid Strychnine

It should be kept in well stoppered glass bottles, as it has a tendency to effloresce in dry air

Solubility—1 in 48 of Water, 1 in 35 of Alcohol (90 p c)

Official in Fr, Mex, Port, Spain and U S

Tests—Strychnine Sulphate when heated to a temperature of $100^\circ C$ ($212^\circ F$) loses its Water of crystallisation, equivalent to 10.59 p c. Anhydrous Strychnine Sulphate melts at $200^\circ C$ ($392^\circ F$). It dissolves in Water, forming a clear solution possessing an extremely bitter taste, and which should be tasted with extreme caution. This solution is neutral in reaction towards Litmus paper. The alkaloid separated by treatment with Ammonia Solution, and in immiscible solvent should answer the tests characteristic of Strychnine given under that heading. The aqueous solution affords with Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. It should yield only a faint yellow coloration when mixed with concentrated Nitric Acid, indicating the absence of Brucine. When ignited with free access of air it should leave no weighable residue, indicating the absence of mineral impurities.

Strychnine Acetate, in colourless, acicular crystals, or as a white crystalline powder, soluble in dilute Acetic Acid. **Strychnine Hydrobromide**, in colourless, translucent, prismatic crystals, or as light, white, silky, acicular crystals, soluble 1 in 65 of Water, 1 in 96 of Alcohol (90 p c), and **Strychnine Valerianate**, in pearly white crystals, or as a white crystalline powder, possessing an odour of Valerianic Acid, slightly soluble in Water, are non official salts of Strychnine, which have in recent years received attention in medical literature

Strychninæ Meta-vanadas has been used in tuberculosis, neurasthenia and atonic dyspepsia.—*B M J E* '01, ii 88

STYRAX PRÆPARATUS.

PREPARED STORAX

FR, STYRAX LIQUIDE PURIFIÉ, GER, STORAX, ITAL, STORACE LIQUIDO,
SPAN, ESTORAQUE LIQUIDO

A light brown, or brownish-yellow semi-liquid, thick balsam, transparent in thin layers, possessing a pleasant aromatic odour, and a sharp pungent balsamic taste. It is obtained from the trunk of *Liquidambar orientalis*. The official product is purified by solution in Ethylic Alcohol, filtration and evaporation of the solvent.

Owing to loss of volatile constituents of the resin during the evaporation of the solvent, Ethylic Alcohol is unsuitable for purification of the resin, and a more volatile solvent would have been preferable, the only objection being greater inflammability. The use of Acetone has been suggested.

It contains free Cinnamic Acid, α and β Storesmol, Styrol and Styriacin (Cinnamyl Cinnamate).

Medicinal Properties—Similar in action to the Balsams of Peru and Tolu. The Ointment (1 to 4) is useful as a parasiticide in scabies and phthieriasis.

Official Preparation—Contained in Tinctura Benzoini Composita.

Not Official—Unguentum Stryacis Compositum, Pommade de Styriax.

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fl., Ger., Hung., Ital., Jap., Mex., Norw., Port., Russ., Span., Swed., Swiss and U.S.

Descriptive Notes—Liquid Storax as imported consists of an opaque greyish viscid liquid, containing about 10 to 20 p.c. of Water, which settles to a certain extent at the bottom of the containing vessel. It is only after purification by solution in Alcohol, filtration and evaporation of the Alcohol, that it presents the appearance required by the *B.P.*, viz., a semi-transparent, brownish-yellow, semi-liquid balsam. It has a strong odour resembling Hyacinth and a balsamic taste. The product of a North American species, *Liquidambar styraciflua*, Linn., is sometimes imported from Guatemala and Nicaragua. It is transparent, of a golden-brown colour, and of the consistence of thick clarified Honey. In Europe it is chiefly used in perfumery; in the United States it is known as Sweet Gum and is used in the preparation of Chewing Gum.

Occasionally the bark of *Liquidambar orientalis*, Mill., from which the Storax has been expressed, is imported under the name of Storax bark, and is utilised in the preparation of tincture, pastilles and incense. The substance sold as Styriax Calamita usually consists of sawdust impregnated with liquid Storax or, more rarely, of the powdered Storax bark 3 parts beaten with Storax 2 parts to cause it to form a mass.

Tests—Storax, according to the *B.P.*, when heated in a test-tube placed in boiling Water becomes more liquid but gives off no moisture. According to Dieterich a limit of moisture might have been introduced into the *B.P.*, and he suggests 15 p.c. as a suitable limit, his former limit of 8 p.c. being deemed too high. When boiled with Potassium Bichromate and Sulphuric Acid an odour of Benzoin is evolved (Essential Oil of Bitter Almonds) is evolved. A soluble or

insoluble in Alcohol might have been included in the *BP*. The *USP* requires that it shall contain not less than 60 p c of its weight of matter soluble in Alcohol (94.9 p c), the alcoholic residue being required to be almost completely soluble in Ether and Carbon Bisulphide, but partially soluble in Petroleum Benzine. The *PG* requires that when 100 parts of Storax are completely exhausted with boiling Alcohol (90 p c) the residue which remains shall amount at the highest to 2.5 p c by weight. The *PG* requires that when mixed with an equal weight of Alcohol (90 p c) it shall form a greyish-brown, cloudy liquid with an acid reaction, which, after filtering and evaporating down, leaves a transparent, semi fluid brown mass, consisting of at least 65 p c by weight of the original Storax, which residue shall be soluble in Ether, Carbon Bisulphide, and Benzol, but not in Petroleum Benzine. Storax leaves 0.01 to 0.5 p c of ash. Useful constants are the Acid and Ester value and the proportion of Cinnamic Acid, but these are not at present included in the *BP*, though it has been recommended that they should be stated. The Acid value varies between 70 to 90, the Ester value from 50 to 120. The *USP* states that Storax when heated in a water bath becomes more fluid, and when agitated with warm Petroleum Benzine the supernatant liquid, on being decanted and allowed to cool, will deposit white crystals of Cinnamic Acid and Cinnamic esters. Agitation with Petroleum Ether has been suggested (*YDP* '01, 116) as a means of detecting Resin as an adulterant. A weighed portion of the Storax mixed with coarse sand is exhausted with Petroleum Ether, the solution filtered, the solvent removed by distillation, and the Saponification value of the residue determined. The Acid value should vary from 40 to 55 and the Saponification value from 180 to 197. Specimens containing Resin as an adulterant may possess an Acid value as high as 116 to 121, and a total Saponification value as low as 172 to 178.

Not Official

UNGUENTUM STYRACIS COMPOSITUM —Oleum Officinale 25, Yellow Wax 15, Liquid Storax 15, Elemi 15, Venetian Turpentine 30 —*Belg*

POMMADE DE STYRAX —Storax 16, Colophony 29, Elemi 16, Yellow Wax 16, Olive Oil 23 —*F*

SUCCI.

JUICES

Juices expressed from fresh medicinal plants, and preserved by the addition of Alcohol, were introduced by Peter Squire in 1835 (*PJ* vol 1). By thus obtaining and preserving the juice of the plant, its properties are not impaired by the action of the heat employed in making an Extract.

Succus Belladonnæ, Succus Conii, Succus Hyoscyami, Succus Scopolari, and Succus Taraxaci, consist of 3 parts of Juice and 1 of Alcohol (90 p c).

Succus Limonis is freshly expressed and contains no Alcohol.

The **Alcoholatures** of the *Fr* are made by digesting equal weights of fresh plant and Rectified Spirit together for 8 or 10 days, pressing and filtering. Aconite, Belladonna, Conium (Ciguë), Digitalis, Eucalyptus, Henbane (*Jus quame*), Stramonium Leaves, Flowers, and Corms of Colchicum, are so prepared.

Not Official
SUCCINUM

AMBER

Translucent or opaque, hard brittle, yellow, yellowish-brown or yellowish-red solid, breaking with lustrous conchoidal fracture

A fossil resinous exudation from *Pinus succinifer*, Goeppert, an extinct coniferous tree, on the shores of the Baltic

Foreign Pharmacopœias—Official in Dutch, Mex (Ambar Amarillo), Port (Ambar), and Swed

OLEUM SUCCINI RECT—A transparent, pale yellow or brownish-yellow limpid oily liquid, possessing a characteristic disagreeable odour and burning acid taste. It is a volatile Oil obtained by the destructive distillation of Amber, and purified by subsequent rectification

A cheaper and inferior product is sold under the name of Oil Succini, which forms a fractional portion of resin spirit obtained by the distillation of ordinary resin—*P J* (4), viii, p 98

Externally it is stimulant and rubefacient

Dose—1 to 3 minims = 0.06 to 0.18 c c

Foreign Pharmacopœias—Official in Hung, Mex, Norw and Port

Tests—Rectified Oil of Amber has a sp gr of about 0.905. It boils between 170° and 186° C (338° and 366.8° F). It has a characteristic unpleasant odour, and a hot acid taste. It is soluble in all proportions of Ether, Chloroform and Carbon Bisulphide

LINIMENTUM SUCCINI—Oil of Amber, 1, Spirit of Camphor, 1, Spirit of Hartshorn, 1

A domestic embrocation for whooping-cough

LINIMENTUM SUCCINI COMPOSITUM—Oil of Amber, $\frac{1}{2}$, Oil of Cloves, $\frac{1}{2}$, Olive Oil, 1—This formula is given in *Pharm Form* as a traditional imitation of Roche's Embrocation, and is now incorporated in the *B P C*

TINCTURA SUCCINI—Amber, 1, Alcohol (90 p c), 16

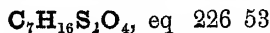
Dose—25 minims = 1.5 c c in Water for headache

Foreign Pharmacopœias—Official in Dutch, 1 Amber and 5, Port, 2.8 Oil in 10, Swed, 1 Amber in 5

SULPHONAL.

SULPHONAL

SULPHONMETHANUM, SULPHONMETHANE



FR, DITHIYLSULFONE-DIMETHYLMETHANE, GER, SULFONAL,
ITAL, SOLFONALE, SPAN, SULFONAL

Colourless, odourless, almost tasteless, prismatic crystals, or a white odourless crystalline powder. Permanent in the air.

The *B P* describes Sulphonal as Dithionophene, a product of the oxidation of Mercaptol obtained from Acetone and Mercaptan. The *U S P* describes it as Dithionophenyl-methylmethane, a product of the condensation of Acetone with Ethyl Mercaptan. The *P G* does not state its origin.

It should be kept in well-stoppered glass bottles.

It belongs to the class of Di sulphones, to which Thional and Tetronal also belong

It is officially required to be in crystals, but it is generally supplied in *powder*, its action is stated to be quicker and more certain in that form than when administered in crystals

Solubility—1 in 500 of Water, 1 in 15 of boiling Water, 1 in 50 of Alcohol (90 p c), 1 in 3 of Chloroform, 1 in 90 of Ether

Medicinal Properties—A pure hypnotic for simple insomnia, when pain is absent. As in repeated doses it may be cumulative, and produce hæmatoporphyrimuria and other toxic effects, it is not the hypnotic to select for continued use. It is more soluble in warm than in cold Water, and still more so in warm alcoholic drinks, the latter being the best way of administration. If taken in the form of a powder or tablet, the action may, owing to comparative insolubility, be deferred for 1 or 2 hours, but if taken in hot solution the action is far more rapid.

Toxic effects following the administration of 35 grains given in divided doses of 20 and 15 grains at an interval of 24 hours. Recovery.—*B M J* '99, i 209

A case in which a quantity of 300 grains was taken in 2 doses of 150 grains, only slight drowsiness supervening owing to prompt action of emetics.—*B M J* '00, i 186

Toxic cumulative effect of Sulphonal and Thional. The only treatment of any avail for Sulphonal poisoning is the free exhibition of alkalis, when there is vomiting and great difficulty in getting the alkali taken in sufficient quantity the intravenous or interstitial transfusion of an alkaline solution might be tried.—*B M J* '92, ii 1250

A fatal case of hæmatoporphyrimuria following its use, also a record of another fatal case in which only 30 grains had been taken in 2 doses.—*B M J* '01, i 1473, *T G* '01, 618

Valuable in early cases of insanity, but in ordinary acute insanity, with extreme restlessness and sleeplessness, Sulphonal even in moderate doses was injurious. The prolonged use of Sulphonal was pernicious from the point of view of auto-intoxication.—*L* '02, i 1539

May be tried for the sleeplessness of uræmic patients, the dose should not be large.—*Pr* lxxvii 658

A serious feature in most of the fatal cases of poisoning is that usually the patients have been under treatment for some time and have been apparently benefited by the drug up to the time of the appearance of toxic symptoms.—*L* '03, i 1023, *B M J* '03, i 853

The urine of patients taking Sulphonal is stated to reduce Fehling's Solution.—*B M J E* '95, ii 43, *P J* (3) xxv 1124

Of the Sulphur hypnotics, Sulphonal was uncertain in its action and caused tissue changes as evidenced by hæmatoporphyrimuria.—*B M J* '05, ii 250

Dose—10 to 30 grains = 0.65 to 2 grammes

Prescribing Notes—It is given in mixtures suspended with Compound Tragacanth Powder 60 grains to 6 fl oz of Water. Also in cachets, capsules, Compressed Tablets, or in powders, to be taken perhaps best of all in hot Water, or hot Spirits and Water

Not Official—Thional and Tetronal

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Fr (Diethylsulfo-nediméthylméthane), Ger, Ital (Solfonale), Jap, Mex, Norw, Russ, Span, Swed, Swiss and U S (Sulphonmethanum)

Tests—Commercial Sulphonal melts at 125.07° C (257.13° F), dried Sulphonal at 125.24° C (257.43° F), purified Sulphonal at

125° 26' C (257° 47' F) The *BP* m.p. is 125° 5' C (258° 11'), this figure is also given in the *USP*. The *PG* states 125° to 126° C (257° to 258° 8' F). At a higher temperature it burns away, emitting an odour of Sulphur Dioxide. When a small quantity is heated with a little powdered Charcoal in a dry tube, or when a mixture of equal weights of Sulphonol and Potassium Cyanide are heated, the characteristic and disagreeable odour of Mercaptan is evolved, if the residue from the latter be dissolved in Water, an excess of Hydrochloric Acid added it yields on the addition of a few drops of Ferric Chloride T.S. a reddish colour. When gradually warmed with dried Sodium Acetate it evolves Hydrogen Sulphide. The saturated aqueous solution should be neutral in reaction towards Litmus paper.

The more generally occurring impurities are Chlorides and Sulphates, readily oxidisable organic impurities, Mercaptan or Mercaptol and mineral impurities. The saturated aqueous solution should yield no turbidity upon the addition of Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of Chlorides and Sulphates. Readily oxidisable organic impurities may be detected by the test with Potassium Permanganate given in the small type below. The substance should leave no weighable residue when heated with free access of air, indicating the absence of mineral impurity, and the solution in boiling Water should be free from odour, indicating the absence of Mercaptan and Mercaptol.

Potassium Permanganate—If 1 drop of Potassium Permanganate be added to an aqueous solution (to 10 c.c. of an aqueous solution 1-50, *PG*), the liquid should not be immediately decolorised, *PG* and *USP*.

Not Official

TRIONAL, Methyl-Sulphonol, Sulphonethylmethanum, Sulphonethylmethane ($C_4H_7S_2O_4$, eq. 240.44)—A white crystalline powder with a faintly bitter taste. Is analogous in composition to Sulphonol, but with a Methyl group replaced by Ethyl.

The *USP* describes Trional under the heading of Sulphonethylmethanum as Diethylsulphonemethylethylmethane, a product of the oxidation of the Mercaptol obtained by the condensation of Methyl ethylketone with Ethyl Mercaptan.

- It should be kept in well-stoppered glass bottles.
- It is not official in the *BP*.

Solubility—1 in 320 of Water, 1 in 11 of Alcohol (45 p.c.)

Medicinal Properties—A pure hypnotic like Sulphonol, but, being more rapidly acting, is more rapidly useful in melancholia, mania, and in many nervous affections, in delirium tremens, in sleeplessness of children, it may induce constipation.

In the insomnia of tricuspid incompetency 10 grains given at bedtime yield successful results.—*MP* '04, ii 515. It requires watching, as cases of artificial nephritis are recorded.

Poisonous effects not produced if used cautiously.—*B M J E* '96, i 27.

In insomnia and delirium due to alcoholism and nicotineism, 10 to 30 grains given at first, after a few trials 15 grains every 4 hours and up to 90 grains per diem were given.—*L* '01, ii 223.

5 to 10 grains thrice daily in chorea.—*B M J* '01, ii 1805, '02, i 267.

15 grains every other night for 15 weeks produced toxic symptoms.—*T* '03, i 1023, *B M J* '03, i 853.

Acute poisoning caused by taking 2½ grains tablets in 1 dose. Recovery.—*L* '03, i 1096.

Dose—For children, 5 to 10 grains = 0.32 to 0.65 gramme, for adults, 15 to 30 grains = 1 to 2 grammes, usually given in cachets followed by a hot drink, or suspended with Tragacanth like Sulphonah.

Ph. Ger. maximum single dose, 2 grammes, maximum daily dose, 4 grammes

Foreign Pharmacopœias—Official in Austria and Spain (Trional), Belg., Dutch, Ger., Jap. (Methylsulfonyl), Fr. (Diethylsulfonyl diethylmethane), Swiss, Diethylsulfonylmethylmethane, U.S. (Sulphonethylmethane)

Tests—Trional melts at 76.5° C (169.7° F). The *U.S.P.* and the *P.G.* give 76° C (168.8° F). At a red heat it evolves an odour of Sulphur Dioxide. When heated with an equal quantity of powdered Charcoal in a dry test tube it evolves the characteristic disagreeable odour of Mercaptan, and when gradually heated with dried Sodium Acetate it emits a characteristic and disagreeable odour of Hydrogen Sulphide. In the directions for performing this test in the case of Sulphonah, the *U.S.P.* directs the use of dry Sodium Acetate, whilst, in the case of Trional, the use of dried Sodium Acetate is directed.

The more generally occurring impurities are Chlorides and Sulphates, readily oxidisable organic impurities, Mercaptan or Mercaptol and mineral impurities. The saturated aqueous solution should not afford a turbidity with Silver Nitrate Solution or with Barium Chloride Solution, indicating the absence of Chlorides and Sulphates. It should not decolorise 1 drop of a 1 in 1000 Potassium Permanganate Solution, indicating the absence of readily oxidisable organic impurities. 1 gramme dissolved in 50 c.c. of boiling Water should evolve no odour, indicating the absence of Mercaptan and Mercaptol. When ignited with free access of air it should leave no weighable residue, indicating the absence of mineral impurities.

TETRONAL (Diethylsulphonyl diethylmethane) - It is analogous in composition to Sulphonah, but with 2 Methyl groups replaced by Ethyl. A white crystalline odourless powder. Soluble 1 in 550 of Water, 1 in 12 of Alcohol (90 p.c.).

It is a hypnotic resembling Sulphonah.

Dose—10 to 20 grains = 0.65 to 1.3 gramme

Tests—Tetronal melts at 85° C (185° F). It is but sparingly soluble in Water, but dissolves in Alcohol, forming a solution which is neutral in reaction towards Litmus paper. When heated with an equal weight of powdered Charcoal in a test tube it evolves the characteristic and unpleasant odour of Mercaptan. When gradually heated with dry Sodium Acetate it evolves the characteristic unpleasant odour of Hydrogen Sulphide. When boiled with 50 times its own weight of Water no unpleasant odour should be developed, indicating the absence of Mercaptan and Mercaptol. The saturated aqueous solution should yield no turbidity on the addition of Silver Nitrate Solution, or on the addition of Barium Chloride Solution, indicating the absence of Chlorides and Sulphates, nor should it immediately decolorise 1 drop of a 1 in 1000 Potassium Permanganate Solution, indicating the absence of readily oxidisable organic impurities.

Not Official SULPHUR

SULPHUR

S, eq. 31.82

Sulphur occurs native, and is found in masses or in the powdery form mixed with various impurities. It is abundant in volcanic countries, as in Sicily, and in some parts of Italy. It readily volatilises, and when the vapours are passed into a large brick chamber kept cold, it condenses in fine powder (Sublimed Sulphur), and when a small chamber is used and kept at a temperature of about 120° C (248° F), it condenses in the liquid form and is run into moulds (Roll Sulphur).

Foreign Pharmacopœias—Official in Belg., Sulphur, Ital. (Solfio), Port. (Enxofre), Mex. and Span., Azufre, Swed.

SULPHUR PRÆCIPITATUM.

PRECIPITATED SULPHUR

B P Syn—MILK OF SULPHUR

A light-grey, or greyish-yellow, smooth, amorphous powder, sometimes possessing a slight odour of Hydrogen Sulphide

It is prepared by precipitating the Sulphur from solution of Calcium Sulphide and Thiosulphate by means of Hydrochloric Acid, the former solution is prepared by boiling Sulphur with Lime

LAC SULPHURIS of former Pharmacopœias contained a large amount of Calcium Sulphate, owing to Sulphuric Acid being used in its preparation, but as Hydrochloric Acid is now employed, no distinction should be made between Milk of Sulphur and Precipitated Sulphur

Medicinal Properties—Similar to those of Sulphur Sublimatum, only more active Mixed with Milk and rubbed till smooth, children take it readily

Dose.—20 to 60 grains = 1 3 to 4 grammes

Official Preparation—Trochiscus Sulphuris

Not Official—Lotio Sulphuris, Trochiscus Sulphuris Compositus, Pastillus Sulphuris Compositus, Sulphur Hair Lotion, Lotio Plumbi et Sulphuris, Unguentum Sulphuris et Plumbi, Unguentum Sulphuris Camphoratum, and Unguentum Sulphuris et Resorcin

Foreign Pharmacopœias—Official in Fr (Soufre Précipité), Ital (Solfo Precipitato), Port (Enxofre Precipitado), Mex and Span (Azufre Precipitado)

Tests—Precipitated Sulphur melts at 115° C (239° F) It should be readily and completely soluble in Carbon Bisulphide When ignited it burns with a blue flame, emitting a penetrating distinctive odour of Sulphur Dioxide When evaporated to dryness with Nitric Acid the residue dissolved in Water and the solution filtered, it yields on the addition of Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid The *B P* requires that under the microscope it shall consist of opaque globules without crystalline matter The *U S P* requires that it shall contain not less than 99.5 p.c of pure Sulphur, but does not indicate a method of determination Neither the *B P* nor the *P G* gives the percentage required

The more generally occurring impurities are acid or alkali, Arsenic, Calcium, Chlorides, Sulphates and mineral impurity The presence of acid or alkali may be detected by the reaction with Litmus given in small type below Arsenic may be detected by the Hydrogen Sulphide test described below If a portion of the specimen be shaken with Water acidified with Nitric Acid and filtered, the filtrate should yield only the faintest turbidity with Silver Nitrate Solution, indicating the absence of more than traces of Chlorides, neither should it yield a turbidity with Ammonium Oxalate Solution When ignited with free access of air it should leave no weighable residue

Litmus—If 5 c.c. of Water be agitated with 2 grammes of Precipitated Sulphur, the liquid should not change the colour of blue or red Litmus paper, *U.S.P.*, moistened with Water, it should not redden blue Litmus paper, *P.G.*

Hydrogen Sulphide—Precipitated Sulphur allowed to stand with 20 parts of Ammonia T.S. at 35° to 40° C (95° to 104° F), with intervals of occasional shaking, gives a filtrate which should not be coloured yellow when acidulated with Hydrochloric Acid or on the subsequent addition of T.S. of Hydrogen Sulphide, *P.G.* If 1 gramme be digested for several hours with 10 c.c. of Ammonia Water and the liquid filtered, one half of the clear filtrate should not leave a residue on evaporation if the remainder be evaporated to dryness on a water bath, then after adding 1 c.c. of Nitric Acid and again evaporating the solution obtained by dissolving the residue in 10 c.c. of Hydrochloric Acid (8 p.c.) should not respond to the modified Grignani test for Arsenic, *U.S.P.*

Preparation

TROCHISCUS SULPHURIS SULPHUR LOZINGI

Contains 5 grains of Precipitated Sulphur, and 1 grain Acid Potassium Tartarate in each, flavoured with Tincture of Orange

Dose—1 to 6 lozenges

Belg. has Sulphuris Tabella, 0.1 gramme = 1½ grains in each, Fr. has Tablettes de Soufre, 0.1 gramme = 1½ grains in each, Mex. has Pastillas de Azufre, 0.1 gramme = 1½ grains in each

Not Official

LOTIO SULPHURIS—Precipitated Sulphur, ½ oz. Glycerin, 120 minims, Alcohol (90 p.c.), 1 fl. oz., Rose Water, 3 fl. oz., Lime Water, 3 fl. oz. Recommended in acne of the face—*L.* '87, 166

This has been incorporated in the *B.P.C.* as follows—Precipitated Sulphur, 6, Glycerin, 3, Alcohol, 12.50, Rose Water, 40, Lime Water, *q.s.* to produce 100

TROCHISCUS SULPHURIS COMPOSITUS—Each lozenge contains 5 grains of Precipitated Sulphur, and 1 grain of Cream of Tartar

These lozenges differ from the official Sulphur lozenge in that they contain no Orange, and are therefore preferred by many

A convenient form of administering Sulphur as a general laxative, in cases of sluggish liver, bleeding piles, and habitual constipation—*L.* '89, 1665

PASTILLUS SULPHURIS COMPOSITUS—Precipitated Sulphur, 5 grains, Acid Potassium Tartarate, 1 grain—*Martindale and B.P.C.*

SULPHUR HAIR LOTION—Acetate of Lead, 1½ drms., Milk of Sulphur, (Calcareaous), 3 drms., Glycerin, 10 drms., Heliotrope Perfume, 2 drms., Water, to 10 oz.—*Pharm. Form.*

Lotio Plumbi et Sulphuris *Syn.* Sulphur Hair Restorer—Lead Acetate, 1.75, Precipitated Sulphur, 3.50, Glycerin, 12.50, Distilled Water, *q.s.* to produce 100—*B.P.C.*

UNGUENTUM SULPHURIS CAMPHORATUM—Precipitated Sulphur, 10 grains, Carbolic Acid, 15 grains, Resorcin, 15 grains, Camphor, 15 grains, Solution of Coal Tar, 25 minims, Benzocated Lard, 240 grains, Soft Paraffin, white, 240 grains—*St. Mary's*

This has been incorporated in the *B.P.C.*

UNGUENTUM SULPHURIS ET RESORCINI—Precipitated Sulphur, 20 grains, Resorcin, 15 grains, Soft Paraffin, yellow, to 1 oz.—*St. Thomas's*

This has been incorporated in the *B.P.C.* as follows—Precipitated Sulphur, 4.50, Resorcin, 3, Soft Paraffin, yellow, to produce 100

UNGUENTUM SULPHURIS PRÆCIPITATI—Precipitated Sulphur, 2, Potassium Carbonate, 1, Lard, 8. Excellent for scabies

SULPHUR SUBLIMATUM.

SUBLIMED SULPHUR

B P Syn—FLOWERS OF SULPHUR

A bright yellow or greenish-yellow amorphous powder, possessing a faint characteristic odour

It may be prepared from native Sulphur or Sulphides

Solubility—Insoluble in Water Slightly soluble in hot Alcohol Only partially soluble in Carbon Bisulphide

Medicinal Properties—Laxative, alterative, diaphoretic, expectorant Employed internally in hæmorrhoids and chronic rheumatism, hepatic congestion, gout, chronic bronchitis and many skin diseases, externally also for skin diseases, especially scabies and acne

Dusted on the membrane in diphtheria—*B M J* '93, ii 993, '94, i 459, *L* '95, i 265, 327 20 grains with or without 5 grains of Dover's Powder 3 times daily in dysentery—*L* '01, ii 1406

In typhoid fever, 20 grains every 2 hours up to 154 grains in the day for adults, for children 5 to $7\frac{1}{2}$ grains every 2 hours up to 60 grains in the day—*B M J E* '02, ii 88

Dose—20 to 60 grains = 1.3 to 4 grammes

Official Preparations—*Confectio Sulphuris* and *Unguentum Sulphuris*, contained in *Pulvis Glycyrrhizæ Compositus* Used in the preparation of *Acidum Sulphuricum*, *Acidum Sulphurosum*, *Emplastrum Ammoniaci cum Hydrargyro*, *Emplastrum Hydragyri*, *Antimonium Sulphuratum*, *Potassa Sulphurata*, *Sulphur Præcipitatum* and *Sulphuris Iodidum*

Not Official—'Chelsea Pensioners,' *Unguentum Sulphuris Compositum*, *Vasolimentum Sulphuris*, *Vasolimentum Sulphuris Compositum*, *Parogenum Sulphuris*, *Parogenum Sulphuris Compositum*

Foreign Pharmacopœias—Official in all Austri (*Sulphur Depuratum*), Belg (*Sulphur Lotum*), Dan (*Sulphur Sublimatum*), also *Sulphur Sublimatum Venale*, Dutch, Ger, Jap and Russ (*Sulphur Sublimatum*, also *Sulphur Depuratum*), Fl (*Soufre Sublime* and *Soufre Sublimé Lave*), Hung (*Sulphur Sublimatum*, also *Sulphur Sublimatum Lotum*), Ital (*Solfo Sublimato*, also *Solfo Sublimato e Lavato*), Mex (*Azufre Sublimado y Lavado*), Norw (*Sulphur Sublimatum*), Port (*Enxofre Sublimado*, also *Enxofre Lavado*), Span (*Azufre Sublimado*, also *Azufre Lavado*), Swed (*Sulphur Sublimatum*, also *Sulphur Sublimatum Elotum*), Swiss (*Sulphur Sublimatum Crudum*, also *Sulphur Lotum*), US (*Sulphur Sublimatum*, also *Sulphur Lotum*)

Tests—Sublimed Sulphur melts at about 115° C (239° F) When ignited it burns with a blue flame, evolving a distinctive penetrating odour of Sulphur Dioxide, which blackens a strip of paper moistened with Mercurous Nitrate Solution When oxidised with Nitric Acid, the residue dissolved in Water yields, with Barium Chloride Solution, a dense white precipitate insoluble in Hydrochloric Acid It is officially required to consist of almost opaque, irregular particles and to be free from admixture of crystalline matter when

examined under the microscope. The *USP* requires that it shall contain not less than 99 p.c. of pure Sulphur, but does not indicate a method for its determination. Neither the *BP* nor *PG* states the requisite percentage nor a method of determination.

The more generally occurring impurities are acid or alkali, Arsenic and mineral matter. If the specimen be shaken with Water and filtered, the filtrate should possess neither an acid nor an alkaline reaction towards Litmus paper, indicating the absence of acid or alkali. The *BP* test of freedom from acidity can only be expected from washed Sulphur, which is official in most Foreign Pharmacopœias. Commercial Sublimed Sulphur is always more or less acid. Sublimed Sulphur always gives an acid reaction unless freshly washed and dried. When shaken with Ammonia Solution and filtered, the filtrate should not, on acidification with Hydrochloric Acid, afford a yellow precipitate or turbidity, nor should another portion yield a residue when evaporated to dryness, indicating the absence of Arsenic Sulphide. The residue left in this Ammonia test might be Ammonium Sulphate, and is no proof of the presence of Arsenic or Arsenic Sulphide. The Arsenic test is not delicate enough. A standard has been suggested (*CD* 08, 1796) of 2 parts per million for Arsenic. When ignited with free access of air Sulphur should burn leaving no weighable residue.

Preparations

CONFECTIO SULPHURIS CONFECTION OF SULPHUR

Sublimed Sulphur, 4 oz., Acid Potassium Tartrate, 1 oz., Tragacanth, in powder, 18 grains, Syrup, 2 fl. oz., Tincture of Orange, $\frac{1}{2}$ fl. oz., Glycerin, $1\frac{1}{2}$ fl. oz. (1 in 2 $\frac{1}{2}$)

Now made with Glycerin, Syrup, and Tincture of Orange in place of Syrup of Orange Peel.

Dose—60 to 120 grains = 4 to 8 grammes

UNGUENTUM SULPHURIS SULPHUR OINTMENT

Sublimed Sulphur, finely sifted, 1, Benzoinated Lard, 9 (1 in 10)

In *BP* '85 it was 1 in 5

Precipitated Sulphur makes a more active Ointment, and Essence of Lemon covers the odour.

An ointment $\frac{1}{2}$ of *BP* '85 strength exerts a destructive effect on the ring worm fungus.—*BMJ* '89, 1398

Foreign Pharmacopœias—Official in Belg. (Sulphuris Alcalini Unguentum), Potassium Carbonate 10, Water 5, Sulphur 20, Lard 65, Jap. and Russ., Sulphur 1, Lard 2, Fr., Sulphur 1, Almond Oil 1, Benzoinated Lard 8, Mex., Sulphur 1, Benzoinated Lard 3, Port. and Swiss., Sulphur 3, Lard 7, Port. has also compound ointment 1 in 5, Russ. has also compound ointment 1 in 10, Span., Sulphur 1, Lard 4, U.S., Sulphur 3, Benzoinated Lard 17.

Not Official

'**CHELSEA PENSIONER**'—Sulphur 6 Mustard, 6 Powdered Guaiacum, 3, Rhubarb, $1\frac{1}{2}$, Nitric, $1\frac{1}{2}$ min. Honey or Treacle sufficient to make it into an Electuary.

Dose—A teaspoonful every alternate night for rheumatism, it is also taken in the morning as an aperient to regulate the bowels

See also GUAIACUM, p 583

UNGUENTUM SULPHURIS COMPOSITUM. *Syn* UNG AD SCABIELEM VIENNENSE WILKINSON'S OINTMENT

Sulphur, 15, Chalk, 10, Tar, 15, Lard, 30, Soap, 30

This has been incorporated in the *B P C*

Formula —Official in Austria, Sulphur 16, Chalk 4, Tar 16, Lard 1 Soap 32, Dutch, Sulphur 15, Pulvis Marmoris 20, Yellow Vaseline 30, Potash Soap 20, Oil of Cade 15, Hung, Sulphur 15, Chalk 10, Potash Soap 30, Lard 20, Yellow Wax 10, Tar 15, Norw and Swed, Sulphur 15, Chalk 10, Tar 15, Lard 30, Soap 30, Swiss, Sulphur 10, Zinc Sulphur 10, Soap 15, Lard 65

UNGUENTUM SULPHURIS COMPOSITUM—Sulphur, 4 oz, Powdered White Hellebore, 10 drms, Nitrate of Potash, 2 scruples, Soft Soap, 4 oz, Lard, 12 oz, all by troy weight—*P L* 1851

VASOLIMENTUM SULPHURIS—Sublimed Sulphur, 3, Linseed Oil, 37, Simple Vasoliment, *q s* to make 100 Heat the Sulphur and Linseed Oil together until dissolved, and make up with Simple Vasoliment—*Y B P* '01, 212

Parogenum Sulphuris *Syn* Sulphur Vasoliment—Sublimed Sulphur 3, Linseed Oil, 37, Parogen, *q s* to produce 100—*B P C*

VASOLIMENTUM SULPHURIS COMPOSITUM—Sulphur Vasoliment, 10, Cade Oil, 10, Thymol, 0 3, Eucalyptol, 3, Turpentine, 30, and make up with Vasoliment to 100—*Y B P* '01, 212

Parogenum Sulphuris Compositum. *Syn* Compound Sulphur Vasoliment—Sulphur Parogen, 10, Oil of Cade, 10, Thymol, 0 3, Eucalyptol, 3, Oil of Turpentine, 30, Parogen, *q s* to produce 100—*B P C*

Not Official

SULPHURIS CHLORIDUM

SULPHUR CHLORIDE

S_2Cl_2 , eq 134 02

A mobile reddish yellow liquid, sp gr 1 69, with a penetrating disagreeable odour, and fuming strongly in air Prepared by the direct union of Chlorine with Sulphur It dissolves without decomposition in Carbon Bisulphide or Benzol, but is decomposed by Water, Alcohol or Ether

UNGUENTUM SULPHURIS HYPOCHLORITIS—Sublimed Sulphur, 1 oz, Sulphur Chloride, 1 fl drms, Spermaceti Ointment (*B P* 1867), 8 oz, Essential Oil of Almonds, 80 minims, is usually added to mask the disagreeable odour

Used in the treatment of scabies and venere

Occasionally made of twice this strength

Sublimed Sulphur, 12, Sulphur Chloride, 2, Essential Oil of Almonds, by weight, 2, Lard, 84—*B P C*

SULPHURIS IODIDUM.

SULPHUR IODIDE

Consists of crystalline masses, possessing a metallic lustre and evolving a strong odour of Iodine It should be kept in well-stoppered bottles in a cool place Like Iodine it stains the skin. It is prepared

by direct combination of Iodine and Sulphur by heating them together

The proportions of Iodine and Sulphur are used in equivalents to form SI_2 , eq 157.72, but the combination is a very loose one

Solubility—1 in 16 of Glycerin, 1 in 4 of Carbon Bisulphide Insoluble in cold Water

Medicinal Properties—The Ointment is an excellent remedy for acne rosacea, and for parasitic, tubercular and other diseases of the skin

Official Preparation—Unguentum Sulphuris Iodidi

Foreign Pharmacopœias—Official in Mex (Yoduro de Azufre), Port (Enxofre Iodado), US (Sulphuris Iodidum)

Tests—Sulphur Iodide when exposed to the air gradually loses Iodine. When heated the Iodine sublimes first. When boiled with Water the Iodine passes off in vapour, the Sulphur remaining as an insoluble residue. The amount of this residue is officially required to be about one-fifth of the weight of the original Sulphur Iodide. The *USP* states that continued boiling with Water vaporises all the Iodide, leaving about 20 p.c. of Sulphur as a residue. It is completely soluble in Carbon Bisulphide. No requisite percentage of pure Sulphur Iodide is mentioned in the *BP*, but a rough method of determination is given based upon the insoluble residue remaining when the Sulphur Iodide is boiled with Water, which is officially required to amount to about one-fifth of the weight of the sample taken. Solution in Potassium Iodide and titration with Sodium Thiosulphate would be a better test for quality than the determination of residual Sulphur. The *USP* requires that it shall contain not less than 70.5 p.c. of Iodine as volumetrically determined by dissolving a mixture of 0.5 of a gramme of the finely-powdered Sulphur Iodide and 1 gramme of Potassium Iodide in 20 c.c. of Water and titrating with Tenth normal Volumetric Sodium Thiosulphate Solution, using Starch Mucilage as an indicator, not less than 28 c.c. should be required, 1 c.c. of Tenth normal Volumetric Sodium Thiosulphate corresponds to 0.01259 gramme of Iodine. This percentage of Iodine corresponds to about 88.3 p.c. of Sulphur Iodide of the formula given above

Preparation

UNGUENTUM SULPHURIS IODIDI SULPHUR IODIDE OINTMENT

Rub 20 grains of Sulphur Iodide with 20 grains of Glycerin to a smooth paste in a warmed mortar, and gradually add 460 grains of Benzoated Lard, and stir until cold

BP (1898) reduced the strength from 1 in 15½ to 1 in 25. Glycerin was added and Benzoated Lard replaced Hard and Soft Paraffin

It is apt to be gritty unless carefully made, it becomes darker on keeping,

SUMBUL RADIX.

SUMBUL ROOT

The dried transverse slices of the Root of *Ferula Sumbul*

Imported from Russia. It possesses a powerful odour resembling Musk.
An inferior kind has of late years replaced the old Sumbul root.

Medicinal Properties—Carminative and antispasmodic, said to be useful in hysteria and allied nervous complaints.

Official Preparation—Tinctura Sumbul

Foreign Pharmacopœias—Official in Mex., Port. (Sombula), U.S. has Extract and Fluid Extract.

Descriptive Notes—The Sumbul Root of commerce is probably not that of the official species, *Ferula Sumbul*, Hook. f., which apparently has not been collected for many years (an undetermined species, *USP*). The segments of the root of *Ferula Sumbul* are 3 to 4 in (75 to 100 mm) in diameter, $1\frac{1}{2}$ to 2 in (37 to 50 mm) in thickness (1 to 3 in (25 to 75 mm) in diameter, $\frac{1}{4}$ to 1 in (18 to 25 mm) or more in thickness *BP*). Externally the bark is papery, pale brown, annulated in the upper or rootstock portion and sometimes bristly with the remains of the leaf-stalks near the apex. The transverse section is spongy and fibrous, and shows an irregular, somewhat contorted arrangement of the vascular tissue, and owing to the exudation of oleoresin it is usually mottled with blackish patches. The root yields about 9 p.c. of soft resin and $\frac{1}{2}$ p.c. of a dingy, bluish, essential oil. It has a bitter and musky taste and a musky odour. The article at present in commerce is probably the root of *Ferula suaveolens*, This has quite a weak musky odour and occurs in smaller pieces, although in general appearance resembling the official kind. Occasionally there has appeared a false Sumbul in commerce, the Indian Sumbul of Perena, which consists of the root of *Dorema Ammoniacum*, scented with tincture of Musk, it closely resembles Sumbul in appearance, but when kept in a bottle or closed vessel the odour of Ammoniacum soon overcomes that of the Musk and is easily recognised, the characteristic taste of Ammoniacum is also easily detected. This root is largely imported into Bombay and is used as incense by the Parsees in their fire temples.

Tests—Sumbul Root contains from 5 to 6 p.c. of ash.

Preparation

TINCTURA SUMBUL TINCTURE OF SUMBUL

Sumbul Root, 1, Alcohol (70 p.c.), 10, by maceration.

Now 1 in 10 instead of 1 in 8, and Alcohol (70 p.c.) used in place of Rectified Spirit.

Dose.— $\frac{1}{2}$ to 1 fl. dr. = 1.8 to 3.6 c.c.

Tests—Tincture of Sumbul has a sp. gr. of 0.895 to 0.900, it contains about 2.5 p.c. w/v of total solids and about 66 p.c. w/v of Absolute Alcohol.

SUPPOSITORIA.

Suppositories are for the most part prepared by the following general formula —

Melt the Oil of Theobroma, triturate the active ingredient intimately with a little of the Oil, and add the mixture to the remainder of the melted Oil in a basin or dish, stir well, and as the mixture begins to thicken pour it into the moulds, which may then be cooled with Water, or in summer by iced Water.

All difficulty in removing the suppositories from the moulds may be obviated by having the moulds previously wiped with oiled Lint.

It is convenient to weigh out ingredients for one or two more suppositories than are required by the prescription. The so called 15 grain moulds, sold for suppositories, do not always hold exactly 15 grains, it is advisable to check their capacity.

In those rare circumstances where moulds are not available, the mixture may be allowed to cool, divided into the requisite number of parts, and shaped into a suitable form.

Hollow cones of suppository shape are made with Oil of Theobroma in various sizes, these can be filled with any desired medicament and closed with a plug, they are known as 'hollow suppositories'.

In India and the Colonies a quantity of Beeswax may be added to suit the temperatures for the time being, so as to produce a desirable consistence.

Cocoa nut Stearin (p 1154), or a mixture of this with Oil of Theobroma, is a better basis for suppositories than Oil of Theobroma in cold weather.

Not Official

SUPRARENAL GLAND

The suprarenal or adrenal bodies or capsules are ductless glands each consisting of two portions which are distinct from a physiological point of view, the cortex and the medulla. A suprarenal body is thus two distinct and independent organs combined with one another. The fresh healthy glands of the ox or sheep are generally used for preparations. The substance causing a rise of blood pressure (Adrenalin) is found in the medulla only, regarding the functions of the relatively large cortical portion of the gland we have no definite information, although some comparative observations indicate that it may have important specific relations to the growth of the body, particularly to the genital organs. The active principle may be boiled without losing its activity, thus allowing the comparatively easy preparation of sterilised solutions, it is, however, prone to absorb Oxygen from the air and to become less active.

Medicinal Properties—A powerful vaso constrictor and cardiac tonic. It is of the greatest value in cases of sudden cardiac failure. An extract of the glands was first used in the treatment of Addison's disease, and numerous early cases are recorded, some showing beneficial effects, and others little or no improvement. The use of the extract has gradually been extended in other directions. It has been used in asthma, anæmia, cyclic albuminuria, and in diabetes mellitus, in exophthalmic goitre, heart disease and capillary hemorrhages, in hay fever, epistaxis, and nasal catarrh, also in asthma. It has been found of great service in ophthalmic work as it lessens congestion and hastens absorption. It is useful in inflammatory conditions of the conjunctiva, and its use is also indicated in pannus, iritis, keratitis and acute dacryocystitis. It controls hæmorrhage in ophthalmic, nasal and obstetric work.

Applied locally it is a powerful astringent and hæmostatic. 1 drop of a 1 in 50,000 aqueous solution of the active principle blanches the normal conjunctiva within 1 minute. Of the dry extract 5 mg per kilo body weight is

sufficient to produce a maximum effect and about $\frac{1}{100}$ of a grain of the active principle is sufficient to produce a distinct effect upon the heart and arteries of an adult man. It has been recommended in Graves' disease, and as an adjunct to the treatment of lupus by the Finsen light. Administration by the mouth has not in some cases been found to be so efficacious as intravenous or subcutaneous injection. For cases of cardiac failure, it is best given intravenously, injections of $\frac{1}{200}$ to $\frac{1}{100}$ grain of Adrenalin being given. Subcutaneous injections of $\frac{1}{1000}$ to $\frac{1}{500}$ grain are recommended in ophthalmic practice to be used immediately before operation.

For introduction into the nose and ear, a 1 in 5000 solution of the active principle is used, or a 5 per cent solution of the Extract. As an ointment 1 of Liquid Extract to 7 of Lanolin Ointment, as a suppository containing 2 or 3 minims of Liquid Extract.

5 to 20 minims of solution given every 6 hours in the treatment of neurotic heart.—*B M J '04*, 1 1009

A case where an unpleasant disturbance of smell, followed within a few hours by a diffuse urticaria extending over the trunk and arms, attended the use of a small swab saturated with Suprarenal Extract applied to the swollen inferior turbinate.—*M P '05*, 11 805 5. to 10, 15 and 20 grains, administered thrice daily in Addison's diagnosis and an early trial of this form of treatment are of great importance.—*L '05*, 11 524

Injections of Adrenal Extract or Epiprenan very useful in severe shock and in poorer cases by a general anæsthetic, but caution is needed if heart is weak from long and exhausting illness.—*B M J E '05*, 11 52

Tests—The powder of desiccated Suprarenal Glands is partially soluble in Water. 0.5 of a gramme macerated with 25 c.c. of Water for a quarter of an hour and filtered yields a filtrate, which gives on the addition of a drop of Ferric Chloride T.S. an emerald-green coloration, the addition of Iodine Solution produces a deep rose red coloration. It should not yield more than 7 per cent of ash. The active principle of the gland is Adrenalin, discovered by Dr. Jokichi Takamine.

Various preparations of the dried gland, of the extract (solid and liquid), and of the active principle have been introduced for medicinal use. The following include the best known preparations—

GLANDULÆ SUPRARENALES (sicc. pulv.)—A dry, light brown or drab amorphous powder. Partially soluble in Water. 1 part represents about 5 of the fresh gland.

Dose—5 grains = 0.32 gramme. Also supplied in tablets containing 0.1 gramme = $1\frac{1}{2}$ grains.

DESICCATED SUPRARENALES (Suprarenal Capsules of the Sheep).—A light, fawn-coloured, or light brownish-yellow powder, 1 grain of the powder representing 8 grains of the fresh Suprarenals.

Dose—1 to 3 grains = 0.06 to 0.18 gramme.

Official in U.S.

Desiccated Suprarenal Gland Tablets.—Each tablet containing 2 grains of the desiccated gland.

TABLET SUPRARENAL GLAND.—Each tablet represents 5 grains of the gland.

EXTRACTUM GLANDULÆ SUPRARENALÆ HÆMOSTATICUM.—A brown, or dark brown, hygroscopic, amorphous powder, readily soluble in Water. It is a very active preparation of the gland.

Dose—1 to 3 grains = 0.06 to 0.2 gramme.

EXTRACTUM GLANDULÆ SUPRARENALÆ LIQUIDUM.—A Liquid Extract, 1 part of which is equal to 1 of fresh gland.

Extractum Suprarenalum Liquidum.—Macerate 100 of Trimmed Suprarenal Glands in 75 of Glycerin for 24 hours, strain and press, and make up the volume to 100 by means of Glycerin and Water in equal parts.—*B P C*

LIQUOR SUPRARENALIS HÆMOSTATICUS—A pale brownish or pinkish-brown liquid, containing the entire active principle of the gland. It has been used with great success as a **spray** in the treatment of hay fever, coryza, etc.

ADRENALIN $C_9H_{13}NO_3$, eq 181.77—It is the active principle of the suprarenal gland, and in constitution partakes of the nature of an alkaloid. It forms a light, white, or almost white, microcrystalline powder, possessing a slightly bitter taste and leaving a feeling of numbness on the tongue. It is very sparingly soluble in cold Water, but dissolves more readily in hot Water. It dissolves readily in diluted Hydrochloric Acid. It is also soluble in Oleic Acid. The salts are mostly non-crystallisable. Solutions of the active principle readily absorb Oxygen from the air and pass into inactive substances. The active principle in the dry form is perfectly stable, it should, however, be kept in small well closed glass phials of a dark amber tint in a cool atmosphere and exposed as little as possible. The form in which it crystallises is dependent upon the condition of the solution from which it is crystallised, and it has been known to exist in the form of prisms, fine needles, rhombic plates, boat or leaf shaped crystals or in the form of wart like crystals.

Foreign Pharmacopœias—Official in Belg. and Fr., not in the others.

Tests—Adrenalin possesses a weak alkaline reaction towards moistened red Litmus paper, and it also has a faintly alkaline reaction towards Phenolphthalein Solution. The aqueous solution, particularly when alkaline, rapidly absorbs Oxygen from the air and passes from a colourless liquid to a pink, red, and eventually brown one. Its aqueous solution may be boiled without decomposition. The highly dilute aqueous solution has a strong action upon the small blood vessels, 1 drop of a 1 in 10,000 solution when instilled into the eye immediately blanches the conjunctiva. It raises the blood pressure, a dose of even about 0.0000005 gramme being sufficient to produce an effect. The diluted aqueous solution affords with Ferric Chloride T.S. an emerald green coloration, with Iodine Solution a beautiful rose pink coloration is produced. On the addition of Potassium or Sodium Hydroxide to the green coloured solution produced by Ferric Chloride T.S., the colour changes from purple to carmine red, and is destroyed when carefully neutralised with diluted acids, reverting to its original colour. Nitric Acid, Potassium Bichromate Solution and Potassium Ferricyanide Solution produce a similar rose pink coloration to that produced by Iodine Solution. Gold Chloride Solution is immediately reduced by the aqueous solution of Adrenalin, a complete separation of the metal taking place. Although alkaloidal in constitution, it does not possess the chemical properties of an alkaloid, and its solutions are not precipitated by the usual alkaloidal precipitants, e.g., Potassium mercuric Iodide (Mayer's) Solution, Iodo Potassium Iodide (Wagner's) Solution, Picric Acid Solution, Tannic Acid Solution, Phospho Molybdic Acid Solution, and Platinum Chloride Solution. A drop or two of Potassium or Sodium Hydroxide Solution produces a reddish brown coloration when added to Adrenalin, or even a very dilute Solution of Adrenalin, and simultaneously a distinctive disagreeable odour, resembling Hydrogen Phosphide, is produced. This test has been suggested (*P. J.* '07, p. 718) as a distinguishing test for Adrenalin, the following method being used in carrying it out—A small quantity of the dry Adrenalin, or a few drops of the 1 in 1000 aqueous solution, is placed in a small porcelain crucible and mixed with 5 to 6 drops of a 10 p.c. Sodium Hydroxide Solution. The solution gradually becomes coloured, and simultaneously a distinctive odour is developed in a few seconds, more or less according to the amount of the Adrenalin present. In the case of a liquid containing substances likely to interfere with the reaction, these must be removed by previous treatment before applying the test. In the case of oily liquids the following method has been suggested (*P. J.* '07, p. 310). Dissolve from 10 to 20 drops of the oily fluid or an equivalent amount of a solid preparation in about 10 times its volume of Ether and shake in a separatory funnel with 10 to 20 drops of Water acidified with Hydrochloric Acid. When the liquids have separated, the lower aqueous layer is transferred to a small capsule or beaker heated on the water bath for a few minutes to remove the Ether and traces of odorous substances. 6 or 8 drops of a 10 p.c. aqueous Sodium Hydroxide Solution

are added and the whole covered with a watch-glass and set aside for 4 or 5 minutes. In the event of Adrenalin being present, the liquid will have acquired a distinct and peculiar odour suggesting Hydrogen Phosphide. Several methods have been advocated for the determination of Adrenalin. It has been suggested that both the reaction with Iodine and that with Ferric Chloride might afford a means of colorimetrically determining its amount, but none of the processes have been found to possess any real practical value.

A synthetic Adrenalin (Suprarenin) has been produced and is now a commercial article. The synthesis of substances allied to Adrenalin has been investigated by Jowett and a record of the researches appears in the *JCS Trans* '05, 967. The chief difference between the synthetic product and the natural appears to be the optical rotation. The natural Adrenalin is laevorotatory, whilst the synthetic Adrenalin, like most synthetic preparations, is optically inactive, and the synthetic preparation is consequently stated not to possess the same physiological activity as the natural product. As the result of experiments recorded (*PJ* '08, 1 626) it has been shown that in respect to rise of blood pressure both substances are exactly alike, and that the optical activity is probably without any influence. A preparation consisting of three-fourths of the dextro-rotatory modification and one-fourth of the laevorotatory is equally as active as the laevorotatory, although the latter has been alleged to be the sole active modification.

Jowett (*JCS Trans* '04, 192) has established the formula $C_9H_{13}NO_3$ for Adrenalin, and the correctness of this work and formula has been confirmed by Continental authorities working on the same subject.

Medicinal Properties.—It possesses the physiological activity of the medullary portion of the gland, but in a very much enhanced degree. It is the strongest haemostatic known. It is in the form of dilute solution of this active principle, and it is now generally used medicinally. As a daily injection of a 1 in 1000 solution in the treatment of laryngeal papillomata (*BMJ* '04, 11 1224), in the treatment of typhoid (*BMJ* '04, 11 1452), as a 1 in 1000 spray in aromatised form in inflammatory affections of the nose and throat (*L* '04, 11 1160), as an injection of 4 to 8 c.c. of a 1 in 1000 solution in tuberculous pleural effusion (*LJ* '04, 11 1008), in the form of an ointment (1 in 1000) made with a base of soft Paraffin and Lanolin it has been used for nose and throat application.—*PJ* '04, 11 967. Adrenalin solutions have frequently been employed in hæmoptysis, but during the latter part of the year (1904) doubts have been entertained of the

value of the drug either hypodermically or by the mouth in hæmoptysis. (*L* '04, 11 1446, *BMJ* '04, 11 1636, '05, 1 68.)

The number of cases of surgical shock in which it has been used are few (*L* '05, 1 849), but there can be no doubt that it affords a very valuable means of treating this condition. When administered intravenously it should be used in very weak solutions. It is rapidly oxidised by the tissues, and its effects are fleeting, so that to be used effectively for raising the blood pressure it is necessary to administer it by continuous intravenous infusion. The strength of the infusion recommended is 1 in 40,000. Serious symptoms have been shown to follow the intravenous injection of 20 minims of a 1 in 1000 solution. If the blood pressure remains low and the patient continues in a dangerous condition of shock, a solution of Adrenalin in physiological salt solution, in the proportion of 1 in 20,000, should be intravenously infused at a rate of about 3 to 5 c.c. per minute.

It has been recommended (*BMJ* '05, 11 125) in the treatment of serous effusions. As much as possible of the fluid is withdrawn by a 2-way trocar and canula, and through the canula still *in situ*, 1 drim of Adrenalin Chloride (1 in 1000) diluted to $\frac{1}{2}$ oz with sterile Water, is introduced by means of an aspirator. The canula is then removed, the wound closed with Wool. In abdominal cases the abdomen is gently manipulated for 5 minutes and a binder firmly applied.

In the treatment of the hæmorrhage of typhoid (*BMJE* '05, 1 48), 20 minims of a 1 in 1000 solution are injected hypodermically every 8 hours until the hæmorrhage is arrested. By the mouth 10 minims may be given every hour.

It has been found useful (*BMJ* '05, 1 700) in lupus erythematosus owing to its tonic effect.

The dose of Adrenalin has been the subject of much controversy

Maximal dose of Adrenalin and analogous preparations of the suprarenal capsules has been established by R Muller (4 J P '05, 238), who recommends that doses 0.00009 gramme should not be exceeded. This quantity may be increased to 0.00015 gramme in cases where the patient is under the influence of an anæsthetic

The new Belgian Pharmacopœia includes Adrenalin and gives tests by which it can be identified. According to the text the 1 in 1000 solution intravenously injected is fatal to a rabbit in a quantity equivalent to less than 0.001 gramme of the active principle per kilogramme of the animal. A dose of even about 0.0000005 gramme raises the blood pressure. This is the first official recognition of the principle of physiological standardisation.

Uses in ophthalmic surgery—*Journal of the Roy Army Med Corps* 08, 1 58

Externally in form of ointment (1 of Chloride to 1000 of base) in neuritis, neuritis and referred pain, applied along course of nerve involved—*T G '07*, 1 298

Good results in diabetes insipidus—*B M J E '07*, 11 12

Prescribing Notes—*When ordered in aqueous or other solution it is usually dissolved in just sufficient diluted Hydrochloric Acid to effect solution. A convenient basis for an oily preparation for a spray is a mixture of 1 part by weight of Absolute Alcohol, to 4 parts by weight of Castor Oil, previously dissolving the Adrenalin as above directed. The strength of the Adrenalin may vary from 1 in 1000 to 1 in 4000, according to the wishes of the prescriber. Oil of Gaultheria and Eucalyptol can be added as antiseptics.*

ADRENALIN CHLORIDE SOLUTION—A transparent, almost colourless liquid, containing 1 part of Adrenalin Chloride, and 5 parts Chlorotone in 1000 parts of Normal Saline Solution.

It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and exposed as little as possible to contact with the air and light. In neutral or faintly alkaline solution Adrenalin is liable to rapidly change in colour, but if the solution be made faintly acid in reaction, the change in colour does not take place with anything like the same rapidity.

For nasal, aurial and ophthalmic use it may be diluted to form 1 in 2000, or 1 in 5000, or a 1 in 10,000 solution.

Dose—5 to 30 minims = 0.3 to 1.8 c c for internal administration.

Liquor Adreninæ Hydrochloricus—Adrenine, 0.10, Chloroform, 0.50, Sodium Chloride, 0.90, Diluted Hydrochloric Acid, 0.25, Distilled Water, q s to produce 100—*B P C*

INSUFFLATIO ADRENALINI COMPOSITA (Squire)—Adrenalin, 1 grain, Boric Acid, in fine powder, 1 oz, Camphor, in fine powder, 1 oz, Cocaine Hydrochloride, 1 grain, Menthol, 40 grains, Eucalyptus Oil, 10 minims, Lycopodium, 2 oz.

COMPOUND SUPRARENALIN SNUFF—Boric Acid, 240 grains, Camphor, 20 grains, Cocaine Hydrochloride, 1 grain, Suprarenalin Powder, 1 grain, Lycopodium, 2 oz, Menthol, 60 grains, Potassium Chlorate (in powder), 120 grains, Oil of Eucalyptus, 10 minims—*Armour's Formulas*

PULVIS SUPRARENALIS COMPOSITUS *Syn* Suprarenal Snuff—Dry Suprarenal, 5, Boric Acid, in powder, 30, Camphor, 1.50, Menthol, in powder, 3, Oil of Eucalyptus, 1, Lycopodium, q s to produce 100—*B P C*

NEBULA ADRENALINI (Squire)—Adrenalin, 0.1, Sodium Chloride, 0.9, Chlorotone, 0.25, Water, to 100.

Nebula Adreninæ—Hydrochloric Solution of Adrenine, 20, Chloroform Water (1 in 200), sufficient to produce 100—*B P C*

NEBULA SUPRARENALIN ET COCAINÆ—Suprarenalin or Adrenalin Solution (1 in 1000), 90 minims, Cocaine Hydrochloride, 9 grains, Distilled Water, 1 fl oz—*Bournemouth Formulary*

Nebula Adreninæ cum Cocaina -Hydrochloric Solution of Adrenine, 20, Cocaine Hydrochloride, 2, Chloroform Water (1 in 200), *q s* to produce 100 —*B P C*

SOLUTIO SUPRARENINI BORICI —Suprarenin, 0 1, Boric Acid, 0 25, Sodium Chloride, 0 9 Thymol, 0 06 Water, to produce 100

A Suprarenin Borate prepared from synthetic **Suprarenin** may be employed in the preparation of the above solution

Liquor Adreninæ Boricus, Boric Solution of Adrenalin —Adrenine, 0 1, Boric Acid, 0 2, Chloroform, 0 5, Distilled Water, *q s* to produce 100 —*B P C*

SUPPOSITORIA ADRENALINI (Squæ) —Adrenalin, $\frac{1}{2}$ grain, Water, 16 grains, Boric Acid, 1 grain, Anhydrous Lanolin, 24 grains, Oil of Theobroma, *q s* to make 480 grains, divide into 32 suppositories

SUPPOSITORIA ADRENALINI ET ÆSCULIN (Squæ) —Adrenalin, $\frac{1}{2}$ grain, Æsculin, 32 grains, Boric Acid, 1 grain, Water, 16 grains, Anhydrous Lanolin, 24 grains, Oil of Theobroma, *q s* to make 480 grains, divide into 32 suppositories,

SUPPOSITORIA SUPRARENALIN —Suprarenalin, $\frac{1}{2}$ grain, Boric Acid, 1 grain, Distilled Water, 15 minims, Anhydrous Lanolin, 50 grains, Cocoa Butter, 400 grains Dissolve the Suprarenalin and Boric Acid in the Water Mix with the Lanolin Add the melted Cocoa Butter, pour into 15-grain moulds when cooling Each contains Suprarenalin $\frac{1}{10}$ grain equal to 16 minims of the 1 in 1000 *Bournemouth Formulary*

Solutio Adreninæ, Adrenine Solution —Adrenine, 0 1, Boric Acid, Water, 3, Wool Fat, 10 —*B P C*

UNGUENTUM ADRENALINI (Squæ) —Adrenalin, 0 1, Diluted Hydrochloric Acid, 0 2, Water, 2, Soft Paraffin, 33, Hydrous Wool Fat, sufficient to produce 100

Unguentum Adreninæ, Adrenine Ointment —Adrenine, 0 1, Boric Acid, 0 2, Water, 3, Hydrous Wool Fat, 50, Soft Paraffin, to produce 100 —*B P C*

An **Unguentum Adreninæ Mitis**, *B P C*, is prepared by diluting 1 of the above Ointment to 5 with Soft Paraffin and perfuming with Otto of Rose (1 to 1000)

Unguentum Adreninæ Album, White Adrenine Ointment —Adrenine, 0 1, Hydrochloric Acid *q s*, Castor Oil, 5, Absolute Alcohol, 2, White Soft Paraffin, to produce 100 —*B P C*

UNGUENTUM SUPRARENALIN ET COCAINÆ —Suprarenalin, $\frac{1}{2}$ grain, Boric Acid, 1 grain, Cocaine Hydrochloride, 5 grains, Distilled Water, 15 minims, Hydrous Lanolin, 250 grains, Vaseline, 250 grains Dissolve the first 3 ingredients in the Water and mix with the Lanolin and Vaseline Contains Suprarenalin, 1 in 1000, Cocaine Hydrochloride, 1 in 100 —*Bournemouth Formulary*

This has been incorporated in the *B P C*

EPINEPHRIN —A white, or greyish-white powder, which is regarded by Abel and Crawford as the active principle of the suprarenal gland

Its chemical constitution has been recently investigated by Dr Jowett —*B M J E* '99, 1 35, *P J* '03, 1 1, '04, 1 247

SUPRARENALIN —A light yellow, stable non-hygroscopic crystalline powder Slightly soluble in cold Water and in Alcohol It is stated to possess all the therapeutic properties of the Suprarenal Capsules

Suprarenalin Solution —A slightly alkaline stable solution, containing 1 of Suprarenalin in 1000

RENAGLANDIN —A light brown syrupy liquid It is stated to be a concentrated and aseptic fluid extract of suprarenal gland Each fl drim is equivalent to 5 grains of the fresh gland

RENALINE — A greyish white crystalline powder, only slightly soluble in cold Water, more readily soluble in warm Water. It gradually darkens in colour when exposed to the air and light. It forms definite salts with the Acids, the chief salt being Hydrochloride. It is also sold in the form of a 1 in 1000 solution and in glass capsules containing 1, 2 and 5 c c of a sterilised solution (1 in 1000, 1 in 2000, or 1 in 10,000).

NEBULA EXTRACTI SUPRARENALIS Suprarenal Extract, 48 grains, Sodium Sulphate, 10 grains, Boiling Distilled Water, to 1 fl oz = 10 p c solution — *Central Throat*

SUPRARENAL OINTMENT — Liquid Extract of Suprarenal Gland, 50 minims, Liquid Paraffin, 2 drms, Hydrous Wool Fat, to 1 oz. It may be scented with Otto of Rose — *Martindale*

Unguentum Suprarenalis — Liquid Extract of Suprarenals, 10, Liquid Paraffin, 25, Hydrous Wool Fat, *q s* to produce 100. This ointment is sometimes perfumed with Otto of Rose — *B P C*

Not Official

SYRUPI

Syrups are apt to ferment or become mouldy when made with too little Sugar, and to crystallise when too concentrated, or when mixed with Acids or Alcohol. There is no uniformity in the method given in *B P* for the 22 Syrups which are official. In 7 of them the final product is directed to be made to a given volume by the addition of Water or of Syrup, and in 3 of them to a given weight. The sp gr is mentioned in 2 of them, Syrupus, and Syrupus Ferri Iodidi. In the case of Syrupus Sennæ and Syrupus Tolutanus, the fluid is made up to a given volume by the addition of Distilled Water before the Sugar is dissolved in it, but in Syrupus Hemidemi and Syrupus Rosæ no such precaution is taken. Syrupus Aurantii and Syrupus Zingiberis are both mixtures of a Tincture with Syrup, but the latter is made up to a definite volume, the former is not.

Not Official

TABACI FOLIA

LEAF TOBACCO

The dried Leaves of the Virginian Tobacco, *Nicotiana Tabacum*, L.
Official in *B P* '85, but now omitted.

When dry they yield about 20 p c of ash, containing a large proportion of Potassium Carbonate.

The Virginian leaf contains about 6 p c of Nicotine, and is one of the strongest varieties of Tobacco.

Medicinal Properties — A powerful depressant, especially affecting the heart and respiration. Smoked, it is sedative and antispasmodic in various cases of asthma. Occasionally used as snuff for its errhine action, increasing the flow of nasal mucus.

It forms the basis of a proprietary article for the relief of neuralgia of the face.

Nicotine is one of the most powerful and rapid poisons known.

Smoke from both tobacco and hay found to be bactericidal to pathogenic bacteria — *L* '07, 1 1220

Tobacco-juice (a strong infusion) is a powerful insecticide, but some preparations for this purpose contain Arsenic in addition to the Tobacco, and in a case that came under our notice, several animals were killed by the Arsenic.

Antidotes — In case Tobacco has been swallowed, an emetic, stimulant, internal and external. Recumbent position, Tannic Acid, Nux Vomica or Strychnine.

Official in Ger, Mex, Port and Span, Folia Nicotiana

Enema Tabaci—Leaf Tobacco, 20 grains, Boiling Water, 8 fl oz—B P 1867, omitted in B P 1885 and 1898, now included in the B P C with a note that it is rarely used

NICOTINA ($C_{10}H_{14}N_2$, eq 160.98)—A nearly colourless, volatile oily liquid, with an acrid, burning taste, inflammable, miscible with Water, Ether, Alcohol, and the fixed Oils. It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and exposed as little as possible to contact with air and light, as it has a tendency to become darker in colour and to resinify. To this alkaloid Tobacco owes its activity. The most easily crystallised salt is the Acid Tartrate. Nicotine is intensely poisonous, and is seldom, if ever, used therapeutically.

The antidotal action of Strychnine, Eserine and the cruciferous plant *Nasturtium officinale*, to Nicotine has been compared, the result being that the expressed juice of *Nasturtium officinale* is claimed (L '05, 1 1596) to be *par excellence* the antidote to Nicotine.

When injected intravenously Nicotine causes (L '05, 1 851) a tremendous increase in the blood pressure. Its effects, however, are transient, the pressure falling to or even below the normal after a few minutes.

Tests—Nicotine has a sp gr of 1.011. It boils at about 250°C (482°F). It possesses a strong alkaline reaction towards red Litmus. It is powerfully laevogyrate. It may be determined in aqueous solution in the absence of other free bases by direct titration with Tenth-normal Volumetric Sulphuric Acid Solution using Methyl Orange Solution as an indicator of neutrality. 1 c.c. of 1 c.c. of 1 c.c. of Volumetric Acid Solution being equivalent to 0.016098 gramme of pure Nicotine. Nicotine is precipitated by the usual alkaloidal reagents, e.g., Potassium Iodide (Mayer's) Solution, Iodo-Potassium Iodide (Wagner's) Solution, etc. The distinctive reactions by which it may be identified are the formation of a yellow amorphous precipitate rapidly becoming crystalline, on the addition of Picric Acid Solution in excess to a solution of Nicotine, or of a Nicotine salt, by the formation of a yellow crystalline precipitate. On the addition of Water, on the addition of Platonic Chloride Solution to a solution of Nicotine in dilute Hydrochloric Acid, by the formation of a crystalline precipitate, on the addition of Mercuric Chloride Solution to an aqueous solution of Nicotine, the precipitate being soluble in diluted Hydrochloric Acid or in Acetic Acid. The precipitate of this precipitate under the microscope is distinctive. Nicotine is volatile in vapour of steam.

NICOTINÆ SALICYLAS (Eudermol)—Colourless, transparent crystals, or a white, crystalline powder, possessing a faint empyreumatic odour. It is soluble in Water and in Alcohol (90 p.c.). It has been introduced as a remedy for scabies, used in the form of a 1 p.c. ointment made with Vaseline or Lanolin—B M J E '99, 11 47, P J '99, 1 227.

Tests—Nicotine Salicylate dissolves readily in Water, forming a clear solution which is faintly acid in reaction towards blue Litmus paper. It answers the tests distinctive of Nicotine given under the heading of Nicotina. A concentrated aqueous solution, when acidified with Diluted Sulphuric Acid, yields a white precipitate soluble in Ether. If the precipitated acid be separated, washed free from mineral acid, and carefully dried, it should possess the m.p. and answer the tests distinctive of Salicylic Acid given under Acidum Salicylicum.

Not Official.

T A B L E L L Æ.

TABLETS

The tablet is one of the most popular forms for the administration of drugs. Tablets of comparatively high finish can be made extemporaneously at the dispensing counter with care in manipulation and due regard to the composition of the drugs to be compressed. Tablets may be machine made when it is required

to turn them out in a polished coherent state, using as little pressure as necessary for that purpose, or in the form of tablet triturates, which are generally moulded by hand

For the compounding of Compressed Tablets, Messrs E White and R A Robinson, Jun., suggested (*C D* '02, 271, 299, *P J* '02, 140, 172) a mixture of Oil of Theobroma 1, and Starch 3 parts, the Oil being melted and the Starch powder stirred in before cooling, of this mixture 1 part is added to each 4 parts of the powder to be compressed, unless much Sugar be present, when more of the powder is required, it is mixed thoroughly without pressure in a mortar, divided into doses, and each dose compressed. This method is excellent for small quantities of tablets, facilitating the compression, and as a white excipient it is an advantage, the mixture of medicament with the Starch Theobroma excipient should be quite cold before beginning the compression, otherwise it has a tendency to adhere to the parts of the machine and cause trouble. Subsequently Messrs E White and H Rodwell found on further experience (*C D* '03, ii 231, *P J* '03, ii 156, 211) that the Theobroma and Starch excipient is not applicable on a large scale by means of machines with an automatic feed arrangement, except in certain cases where the mixture of substance and excipient happens to form a fairly granular powder capable of flowing evenly and uniformly from the hopper to the die. They have experimented with the object of removing that defect and to devise a method by which, when the tablets are crushed between the fingers or between paper, a soft and smooth powder could be produced. The problem resolved itself into the possibility of devising methods by which the Oil of Theobroma could be uniformly distributed throughout the material to be compressed, forming at the same time a granulated product capable of automatic feeding and compression into a coherent polished tablet with the minimum of force. They recommended the two methods here given.

METHOD I—Theobroma Emulsion

Oil of Theobroma 25, Hard Soap 5, Tragacanth 0.5, Benzoic Acid 0.25, Water to 100. Dissolve the Soap in 25 parts of Water by heat, add the hot solution to the melted Theobroma, and mix by whisking or agitation, shake in the Tragacanth, add the Benzoic Acid, then the remainder of the Water.

Gum Acacia may be used in place of Soap without making any appreciable difference in the general utility of the product. The product in either case should be a thick, smooth, white cream, free from lumps. The addition of Benzoic Acid is only necessary as an antiseptic precaution if the product be kept in stock.

The method of application is as follows—The substance to be compressed, in the finest possible powder, should be triturated with sufficient of the emulsion to form a damp coherent powder—just so damp that it can be shaken through a No 20 or 30 sieve without pressure and without adhering to the meshes. The sifted product, after exposure to the air for a few hours, or during the night, is ready for compression. If the drying process be accelerated by the aid of artificial heat, the dried product must be allowed to stand for an hour or two at least for the Theobroma to solidify before compression is attempted, but in the majority of cases it is better to avoid the use of artificial heat.

If the bulk of substance to be compressed in each tablet either demands or allows the addition of any diluting material, cane sugar is best, in no case does it interfere with the production of a good tablet. When the substance to be compressed is of a dusty nature, and has little tendency to cohere on compression, the addition of a little Glucose is advantageous, the tablet having a better finish and less liability to crack after compression.

METHOD II—Ether-Alcohol Solution of Theobroma.

Oil of Theobroma 1 oz., Ether to 6 oz. Dissolve and add an equal volume of Rectified Spirit as required for use.

The manner of granulating with the above is to add it to the substance or mixture contained in a mortar, trituration being accomplished as quickly as possible, the whole of the solution required being added at once. The mass is then passed through a No 20 or 30 sieve and allowed to dry by exposure

Compression can, in some cases, be proceeded with almost immediately, but it will be found more satisfactory generally to allow the mixture to stand for an hour or two. Sugar granulates remarkably well with the above excipient, and the previous remarks on its addition apply here as well.

Not Official

TALC

CRÆTA GALLICA, FRENCH CHALK, SOAPSTONE

A white, or almost white, impalpable powder, or in greyish irregular masses, possessing a waxy lustre. It has a characteristic saponaceous feeling to the skin, is practically odourless and tasteless. It is a hydrated Magnesium Silicate. It is insoluble in Water, insoluble in dilute acids, and in dilute solutions of alkali Hydroxides.

Foreign Pharmacopœias—Official in F, Ger, Swiss and U.S.

Tests—Talc has a sp gr of about 2.5. When fused with a mixture of anhydrous Sodium and Potassium Carbonates it leaves a residue which, dissolved in hot Water, filtered, the filtrate acidified with Hydrochloric Acid, evaporated to dryness, reacidified with Hydrochloric Acid, again evaporated to dryness, treated with Water and filtered, leaves on the filter an insoluble residue of Silica, when sufficient Ammonium Chloride is added to the filtrate to hold the Magnesium in solution, it yields on the addition of Ammonia Solution a white gelatinous precipitate, indicating the presence of Aluminium, and if the precipitate be removed by filtration, the filtrate affords on the addition of a Phosphate Solution a white crystalline precipitate, indicating the presence of Magnesium. The Talc should not contain more than 5 p.c. of moisture. Diluted Hydrochloric Acid, as determined by boiling a weighed quantity of 1 gramme for 30 minutes with 25 c.c. of Diluted Hydrochloric Acid, maintaining the volume by the addition of Water from time to time, filtering and evaporating the filtrate to dryness, igniting and rapidly weighing.

PURIFIED TALC—A white, or almost white, inodorous, impalpable powder, insoluble in Water, insoluble in dilute mineral acids and insoluble in dilute solutions of the alkali Hydroxides. It is obtained from natural Talc by removing the matter soluble in Hydrochloric Acid by repeatedly boiling a mixture of Hydrochloric Acid and Water, the purified product being washed with Water until a portion of the wash Water is neutral to Litmus, and yields no opalescence with Silver Nitrate Solution, after acidification with Nitric Acid. It is dried at 110° C (230° F).

Medicinal Properties—It is employed as a soothing and protecting powder to the skin and is an ingredient in many 'dusting powders' and 'face powders'. It is also employed as a filtering medium to clarify turbid liquids.

Foreign Pharmacopœias—Official in the U.S.P. It also appears in the B.P. Appendix.

Tests—Purified Talc, when fused with a mixture of anhydrous Sodium and Potassium Carbonates, should answer the corresponding tests given under Talc. The soluble matter should not amount to more than 0.05 p.c. as determined by boiling 10 gms. of the purified Talc for 30 minutes with 50 c.c. of Water, maintaining the volume by the addition of Water from time to time, filtering and evaporating one half the filtrate to dryness. The remaining half, when acidified with Hydrochloric Acid, should yield no blue colour on the addition of Potassium Ferrocyanide Solution, indicating the absence of Iron. When ignited at a dull red heat it should leave a residue amounting to not less than 95 p.c.

BORATED TALC—Boric Acid, in fine powder, 10, Purified Talc 90,

TAMARINDUS.

TAMARINDS

FR, TAMARIN, GER, TAMARINDENKUS, ITAL, TAMARINDO, SPAN, TAMARINDO

The Fruits of *Tamarindus Indica*, L., freed from the brittle outer part of the pericarp and preserved with Sugar

Imported from the West Indies

Medicinal Properties —Refrigerant and slightly laxative In fused with Water, forms a cooling drink in febrile affections, it may also be given with Milk to form **Tamarind Whey** (1 Pulp in 40)

Dose — $\frac{1}{4}$ oz = 7 1 grammes and upwards

Official Preparation —Contained in Confectio Sennæ

Official in all the Foreign Pharmacopœias except Dan, Fr and Ger (a crude and a stained)

Descriptive Notes —The Tamarinds of commerce consist of the fruit deprived of its hard epicarp, and are imported in three forms, viz, West Indian, preserved in Syrup and packed in barrels, East Indian, deprived of the epicarp and pressed into loose masses, and Egyptian, pressed into hard circular flattened cakes, 4 to 8 in (10 to 20 cm) in diameter and 1 to 2 in (25 to 50 mm) thick The pulpy part consists of the mesocarp, the leathery endocarp encloses the seeds Judging from the official description, the West Indian Tamarinds are apparently intended to be used, the pulp should not yield any characteristic reaction for Copper with the test for that metal, which is only likely to be present in the West Indian Tamarinds The cheaper Egyptian Tamarinds are said to be used for curries and sauces, and in the manufacture of tobacco On the Continent East Indian Tamarinds are the kind principally used in pharmacy

Tests —Tamarinds contain an amount of acid equal to about 10 p.c. calculated as Tartaric Acid In the event of Copper vessels being used, Tamarinds are liable to take up this metal

TARAXACI RADIX.

TARAXACUM ROOT

FR, PISSE NELLE, GER, LOWENZAHN, ITAL, TARASSACO
SPAN, HOJA DE TARAXAGON

The fresh and the dried Roots of *Taraxacum officinale*, Wiggers

It is officially required to be collected in the autumn, but the root is best in the very late autumn or winter months, or in the early spring

Medicinal Properties —A mild laxative and bitter tonic, given in atonic dyspepsia with habitual constipation

Official Preparations —Extractum Taraxaci, Extractum Taraxaci Liquidum, and Succus Taraxaci

Not Official —Decoctum Taraxaci, Elixir Taraxaci Compositum, Liquor Taraxaci

Foreign Pharmacopœias —Official in all except Belg, Dan, Dutch and Norw, Fr (Pissenlit), Ital (Tarassaco), Mex (Diente de Leon)

Descriptive Notes.—Taraxacum or Dandelion Root varies in size according to age, from 8 to 12 in (20 to 30 cm) long, and from $\frac{1}{2}$ to 1 in (12.5 to 25 mm) in diameter, and is sometimes branched in the upper portion, due to the original crown of the root being drawn into the earth and giving off lateral buds which form rootstocks. Externally the root is pale brown when fresh, but darker brown when dry, with a short fracture showing a thick white cortex, having numerous translucent concentric rings containing laticiferous vessels, and a yellow porous woody centre.

The root, both fresh and dried, is official, and is directed to be collected in the autumn. The *PG* directs the whole plant to be collected in spring before flowering. The juice of the root quickly undergoes alteration on exposure to the air. The dried root is much attacked by insects, and should not be kept more than a year. The roasted root is used to form Dandelion Coffee.

Tests —Taraxacum Root contains from 4 to 5 p.c. of ash.

Preparations

EXTRACTUM TARAXACI. EXTRACT OF TARAXACUM

Crush fresh Taraxacum Root, press out the juice, allow the feculence to subside, heat the liquid to 212° F (100° C), and maintain the temperature for 10 minutes, strain, evaporate to the consistency of a soft extract.

Dose —5 to 15 grains = 0.32 to 1 gramme

Official in Ital and U.S., from fresh root, Fr., from dried leaves, Austr., Dutch, Hung., Port., Russ. and Swed., from whole plant, Ger. and Jap., from dried root, Mex., from root and leaves.

EXTRACTUM TARAXACI LIQUIDUM. LIQUID EXTRACT OF TARAXACUM

Macerate 20 of dried Taraxacum Root (in No. 20 powder) in 40 of Alcohol (60 p.c.) for 48 hours, press out 10 of liquid, add to the pressed residue 40 of Distilled Water, macerate for 48 hours, press out the liquid, strain and evaporate to 10, mix this with the former 10 to make the total measure 20, filter.

When made in this way it deposits greatly. A much better Fluid Extract is made by percolation with Alcohol (80 p.c.)

Dose.— $\frac{1}{2}$ to 2 fl. drm. = 1.8 to 7.1 c.c.

Foreign Pharmacopœias.—Official in Russ. and U.S.

Tests —Liquid Extract of Taraxacum has a sp. gr. of 1.040 to 1.060, it contains from 16 to 25 p.c. w/v of total solids and about 25 p.c. w/v of Absolute Alcohol.

SUCCUS TARAXACI. JUICE OF TARAXACUM

3 of the expressed Juice from bruised fresh Taraxacum Root, mixed with 1 of Alcohol (90 p.c.), after 7 days, filter.

Dose.—1 to 2 fl. drm. = 3.6 to 7.1 c.c.

Not Official

DECOCTUM TARAXACI—Dried Dandelion Root, 1, Distilled Water, *q s* to produce 20, after boiling for 10 minutes and straining—*B P* 1885 ^α
This has been incorporated in the *B P C*

ELIXIR TARAXACI COMPOSITUM—Fluid Extract of Taraxacum 3 5, Fluid Extract of Wild Cherry 2, Fluid Extract of Licorice 6, Tincture of Sweet Orange Peel 6, Tincture of Cinnamon 3 5, Compound Tincture of Cardamom 3, Aromatic Elixir, *q s* to produce 100 Mix them, allow to stand several days if convenient, and filter. Average dose—8 c c (2 fl drm)—*U S N F* 1896

This has been incorporated in the *B P C*, but *U S N F* 1906 has altered the quantity of Tincture of Cinnamon from 3 5 to 3 0

LIQUOR TARAXACI—A preparation resembling the Succus, but in which the Alcohol (90 p c) is added directly to the bruised root before pressing. Introduced many years before the Succus and superior to it. The opinion (*C D* '92 1 612) is wrong that Liquor in this case is synonymous with Fluid Extract, since the root depreciates considerably in the drying, before powdering

TEREBENUM.**TEREBENE**

A transparent, colourless, mobile, optically inactive liquid

It consists for the most part of the hydrocarbons Dipentene and Terpinene, with some Cymol and Camphene

Terebene is described by the *B P* as a mixture of Dipentene and other hydrocarbons obtained by agitating Turpentine Oil with successive quantities of Sulphuric Acid until it no longer rotates the plane of a ray of polarised light subsequently distilling in a current of steam, the *U S P* describes it as a liquid consisting of Dipentene and other hydrocarbons obtained by the action of concentrated Sulphuric Acid on Turpentine Oil and subsequent rectification with steam

It should be kept in well stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the light

Solubility—1 in 6½ of Alcohol (90 p c), in all proportions of Absolute Alcohol or Chloroform, 1 in 3¼ of Ether, 5 in 8 of Glacial Acetic Acid, very sparingly in Water

Medicinal Properties—Antiseptic A stimulating, disinfecting, expectorant in winter cough (chronic bronchitis) It can be used as an inhalation, mixed with Magnesium Carbonate and hot Water, or from an antiseptic respirator—*B M J* '86, 1 259, 392, '87, 1 796, *P J* (3) xvi 611 In phthisis, *P* lxx 275

Dose—5 to 15 minims = 0 3 to 0 9 c c

Prescribing Notes—*Small doses may be taken on sugar It may be given in mixture suspended with Mucilage of Gum Acacia, in flexible capsules, lozenges or pastils*

Not Official—Vapor Terebenæ, Terpin Hydrate, and Terpinol

Foreign Pharmacopœias—Official in Russ and U S

Tests—Terebene has a sp gr of 0 862 to 0 876, the *B P* states 0 862 to 0 866, the *U S P* states from 0 860 to 0 865 at 25° C (77° F) It is officially stated not to rotate the plane of a ray of polarised light It boils between 165° and 175° C (329° and

347° F) The *BP* states that it should distil between 156° and 180° C (312° 8' and 356° F), but these limits are generally considered too wide, and admit an optically active specimen. Most commercial samples possess a slight action on polarised light. Naylor (*CD* '99, 11 230) could not imagine why optically inactive Terebene was introduced into the *BP*, when the reputation was made on an optically active preparation. *USP* states that it boils at 160° to 170° C (320° to 338° F).

The more generally occurring impurities are acid, undecomposed Turpentine Oil and resinous substances. A piece of blue Litmus paper moistened with Water should not be reddened by a drop of the specimen indicating the absence of acidity. The sample should be almost completely inactive towards polarised light, indicating the absence of undecomposed oil. When evaporated in a porcelain dish on a water-bath, not more than a slight residue should be left, indicating the absence of more than traces of resinous substances. The *BP* requires that not more than 15 p.c. should distil below 165° C (329° F), but this statement requires modification, as it would admit specimens of a very bad quality. It is officially required to leave after distillation only a slight viscid residue, indicating the absence of excess of Resin.

Not Official

VAPOR TERE BENÆ—Pure Terebene, 40 minims, Light Magnesium Carbonate, 20 grains, Distilled Water, to 1 oz.—*Throat and Central Throat*

TERPIN HYDRATE ($C_{10}H_{20}O_2$, H_2O , eq 188.74)—Crystals, glistening, rhombic prisms, or a crystalline powder, possessing a faint, odour, and a somewhat bitter taste.

The *USP* describes it as the Hydrate of the diatomic Alcohol, Terpin. It is official in *USP* and *PG*, but not in the *BP*.

It should be kept in well-closed bottles of a dark amber tint.

Solubility—1 in 280 of Water, 1 in 14 of Alcohol (90 p.c.), 1 in 46 of Alcohol (60 p.c.).

The solubility figures for Terpin Hydrate in Water, Alcohol (90 p.c.) and Alcohol (60 p.c.) given in the *BPC* have evidently been derived from the *Companion*. The figures for the solubility in boiling Alcohol, in Ether, and in Chloroform appear to have been taken from the *USP*, those for the two latter are incorrect for the solubility of the substance in Ether and in Chloroform at the temperatures at which (in the preface) the *BPC* solubilities are stated to have been determined, but are correct for a temperature of 25° C (77° F), provided in the case of Ether, that Ether *USP* (sp. gr. 0.720) is used as a solvent, the point appears to have been ignored in the *BPC* that the Ether *BP* is not Ether *USP*.

Used as an expectorant to reduce secretion in bronchitis and other respiratory disorders.—*Ph*, liv 383.

Dose—3 to 10 grains = 0.2 to 0.65 gramme.

Foreign Pharmacopœias—Official in Dutch, Fr., Ger., Ital., Jap., Mex., Norw., Russ., Span. (Terpina), Swed., Swiss and U.S.

Tests—Terpin Hydrate melts at about 116° C (240.8° F), the *USP* states at 116° to 117° C (240.8° to 242.6° F), when quickly heated, the *PG* 116° C (240.8° F), and loses Water, the m.p. reverting to 102° C (215.6° F). When heated it loses its Water, and at a temperature of 258° C (496.4° F) anhydrous Terpin distils over, solidifying to a crystalline mass possessing a m.p. of about 102° C (215.6° F). It affords an orange yellow colour on treatment with Sulphuric Acid. The hot aqueous solution is rendered

turbid by the addition of a few drops of Sulphuric Acid, a characteristic powerful aromatic odour being simultaneously evolved. It should possess no pronounced terebinthinate odour. It dissolves readily in hot Alcohol (90 p c), to form a clear solution which should not possess an acid reaction towards blue Litmus paper.

TERPINOL—A colourless, or nearly colourless liquid, possessing a strong hyacinthine odour. It is a mixture of Terpenes, with variable proportions of Terpeneol. It has a tendency to thicken and darken on exposure to air and light. It is practically insoluble in Water, but soluble in Alcohol (90 p c) and in Ether.

Official in Spain

Dose—2 minims = 0.12 c c

TEREBINTHINA CANADENSIS.

CANADA TURPENTINE

B P Syn—CANADA BALSAM

A clear, pale yellow, or greenish-yellow, slightly fluorescent, viscous oleo resin, possessing a terebinthinate odour and a somewhat bitter taste.

The oleo resin official in the *B P* is obtained from *Abies balsamea*, Mill. The liquid oleo resin official in the *U S P* is obtained from the same tree. It is also derived from *Pinus Fraseri*, Pursh, in Pennsylvania and Virginia, and from *Abies Canadensis*, Mich.

A solution of the hard brittle solid left on the evaporation of the volatile Turpentine when dissolved in Benzol, Toluol, or Xylol is much used as a medium for mounting microscopical objects, and as a cement for glass. It is also used in its natural state for the same purpose.

Solubility—Soluble in all proportions of Benzol, Chloroform and Ether, 1 in 3 (or less) of absolute Alcohol, 1 in 1 (or less) of Alcohol (90 p c).

Seldom used internally, its medicinal properties are similar to those of Oleum Terebinthinae.

It is used in the preparation of Collodium Flexible.

Foreign Pharmacopœias—Official in U S.

Tests—Canada Balsam by long exposure to the air or quickly when heated, loses about 25 p c of its weight of volatile Oil and forms a hard, brittle solid, which dissolves in Benzol, Toluol or Xylol. It solidifies when mixed with about one sixth of its weight of Magnesia moistened with a little Water, the *U S P* mentions when mixed with 20 p c of its weight of Magnesium Oxide previously moistened with Water. The Ester value of the Balsam varies from 4.5 to 9.8, the Acid value from 84.9 to 85.9, and the Saponification value from 89.4 to 95.7. The Balsam is stated (*U D '04*, 1439) to have a sp gr of 0.987 to 0.994, an optical rotation of $+1^{\circ}$ to $+4^{\circ}$ in a 100 mm tube, a refractive index at 20°C (68°F) of 1.518 to 1.521 and an Acid value of 84 to 87. The Volatile Oil is stated to have a sp gr of 0.862 to 0.865, an optical rotation of -26° to -29° in a 100 mm tube, a refractive index at 20°C (68°F) of 1.472 to 1.477 and an Ester content calculated as Boonyl Acetate of 0.4 to 0.6 p c. The *B P Coder* stated that 'the Volatile Oil consists

chiefly of Lævo-pinene, that the sp gr of the Oil is about 0.987 to 0.994, optical rotation, $+1^{\circ}$ to $+4^{\circ}$, refractive index, 1.518 to 1.521, Acid value, 84 to 87. The word 'Oil' was subsequently altered to 'Turpentine' in the list of additions and collections.

Not Official

TEREBINTHINA CHIA

CHIAN TURPENTINE

An oleo-resin obtained from the incised trunk of *Pistacia Terebinthus*, collected in Scio. A soft solid with a characteristic odour. When treated with its own weight of Absolute Alcohol or pure Ether, the greater portion is dissolved. Was recommended in cancer. — *L* '87, 11 1005, 1144, 1190, 1244

Dose — 5 to 10 grains = 0.32 to 0.65 gramme

Official in Port

PILULA TEREBINTHINÆ CHIÆ — Chian Turpentine, 6 grains, Sublimed Sulphur, 4 grains. To be made into 2 pills, and taken every 4 hours.

A case is reported of these pills forming a compact mass in the bowel, removed by enemas. — *C D* '90, 11 75.

TEREBINTHINÆ OLEUM.

OIL OF TURPENTINE

FR, ESSENCE DE TEREBINTHINE OFFICINALE, GER, TERPENTINOL, ITAL, ESSENZA DI TREMENTINA, SPAN, ESENCIA DE TREMENTINA

A transparent, colourless, or nearly colourless, limpid liquid

The volatile Oil official in the *BP* is obtained from *Pinus sylvestris* and other species of *Pinus*, and is rectified if necessary. The *USP* includes both an Oil of Turpentine and a rectified Oil of Turpentine. The Oil is described as a volatile Oil recently distilled from Turpentine, Turpentine *USP* is described as a concrete oleo-resin obtained from *Pinus palustris*, Miller, and from other species of *Pinus*. The *PG* also includes a Turpentine Oil and a rectified Oil of Turpentine. The Oil of Turpentine *PG* is described as a volatile Oil obtained from different species of *Pinus*. The rectified Oil of Turpentine of the *USP* is prepared by rectifying Turpentine Oil with Sodium Hydroxide Solution (about 5 per cent). That of the *PG* by treatment with Calcium Hydroxide Solution and redistillation, in each case three-fourths of the distillate is collected.

It should be kept in well-closed glass vessels, preferably of a dark amber tint and in a cool atmosphere.

The Oil of Turpentine sold in Britain is almost wholly imported from America, and is the product (mainly) of *Pinus palustris*, Mill., and *P. Teda*, L. German and Russian Oils are principally distilled from *P. sylvestris*, L., French Oil from *P. Pinaster*, Sol. Hungarian Oil of Turpentine is distilled from the cones of *P. umilis*, Haenke, and Carpathian Oil of Turpentine, also known as Riga Balsam, from *P. Cembra*, L.

Oil of Turpentine, when exposed to the continuous action of atmospheric air in presence of Water, develops a large quantity of Hydrogen Peroxide, Camphoric Acid, and other oxygenated products, which form the basis of the 'Sanitas' series of disinfectants.

Oil of Turpentine dissolves Beeswax, Iodine, Sulphur, Phosphorus, fixed Oils, also Resins, forming varnishes.

Solubility—1 in 6½ of Alcohol (90 p c), in all proportions of Absolute Alcohol, Carbon Bisulphide, Chloroform, Ether, sp gr 0.720, and Glacial Acetic Acid

Medicinal Properties—Antiseptic, expectorant, hæmostatic, diuretic, anthelmintic. Useful in passive hæmorrhage from the various organs, 4 fl drms along with an equal quantity of Castor Oil is often successful in removing tapeworm. Antispasmodic in hysterical affections and in hiccough, it is said to dissolve gallstones. In small doses (2 to 10 minims), and in large doses (3 to 4 fl drms), it does not usually tend to irritate the kidneys, but in doses of about 1 fl drim it is apt to do so. Contra-indicated in Bright's disease. Used as an **inhalation** in chronic bronchitis and other lung diseases, as an **enema** with Castor Oil for obstinate constipation, for flatulency and tympanitic distension of the bowels, and in thread-worm. Externally rubefacient and counter irritant, employed as a **liniment** in chronic inflammatory pain and rheumatism, and as a fomentation in acute pain.

10 minim capsules every 2 or 3 hours, or in the form of an **emulsion** with equal parts of Spirit of Chloroform and Spirit of Nitrous Ether, have given good results in enteric fever, but should not be given in albuminuric vesical catarrh.—*B M J* '04, 11 1450. An **enema** of Soap and Water containing 1 oz of Turpentine is of great value where there is flatulent distension of the colon.—*B M J* '04, 11 1452. Its use is stated to check bleeding sometimes, but to be more effectual in melæna than in hæmoptysis.—*B M J* '05, 1 68. Its value as a styptic in typhoid has been questioned, but in the absence of a better remedy it should be used.—*B M J* '05, 1 414.

In hæmoptysis in 10-minim doses in capsules.—*Edin Med Jour* '05, p 467.

In renal hydatids, 15 minims mixed with Liquor Potassæ, Mucilage and Liquorice, night and morning.—*L* '05, 11 601.

Flies and gnats are kept away by the odour of Turpentine.

Dose—2 to 10 minims = 0.12 to 0.6 cc, as an anthelmintic, 3 to 4 fl drms = 10.6 to 14.2 cc.

Prescribing Notes—Usually given in the form of mixture suspended with Mucilage or Powder of Gum Acacia. It may be given in Mistura Amygdalæ. It is also given in capsules. 1 fl drim of Mucilage, with diligent trituration, renders ½ fl drim of Oil of Turpentine emulsive with 1 fl oz of Distilled Water.

30 grains Powder of Gum Acacia rubbed first with 1 fl drim of Oil of Turpentine, then with 1 fl drim of Water, and lastly triturated whilst adding gradually 1 fl oz Distilled Water, makes a good emulsion.

Official Preparations—Linimentum Terebinthinæ and Linimentum Terebinthinæ Aceticum. Used in the preparation of Terebenum.

Not Official—Confectio Terebinthinæ, Emulsum Olei Terebinthinæ, Enema Terebinthinæ, Linimentum Terebinthinæ, Unguentum Terebinthinæ, Vasolimentum Terebinthinæ, and Parogenum Terebinthinæ.

Antidotes—Emetics, Epsom Salts, demulcent drinks, Morphine or Laudanum to relieve pain.

Foreign Pharmacopœias—Official in Austr, Belg, Dutch, Fr, Ger, Hung, Ital, Jap, Norw, Port, Russ, Span, Swed, Swiss and U S.

Austr, Ger, Jap, Swiss and U S have also Rectificatum, Dutch has also Depuratum.

Tests—Rectified Oil of Turpentine has a sp gr of 0.860 to 0.880, the *B.P.* does not give a sp gr, the *USP* states 0.860 to 0.870 at 25° C (77° F), the *P.G.* 0.860 to 0.870. It boils at

about 156° C (312.8° F), which is the figure given in the *BP*. The *PG* states that it distils completely between 155° and 162° C (311° and 323.6° F). The *BP* states that it should distil almost entirely below 180° C (356° F). This temperature is considered (*CD* '98, 11 55) to be too high, boiling at about 155° C (311° F) and at least 80 p.c. distilling below 165° C (329° F) would have been better. The *USP* requires that the larger part of the Oil should pass over between 155° and 162° C (311° and 323.6° F). The optical rotation of the Oil may be either dextrogyrate or laevogyrate. French Oil of Turpentine is strongly laevogyrate (−20° to −40° in a tube of 100 mm length). American Oil of Turpentine is dextrogyrate, the rotation usually varying from +9° to +14°. A 62 lb quantity when fractionally distilled (*CD* '00, 11 174) yielded up to 162.5° C (324.5° F) a distillate (91.2 p.c. of the whole) which was entirely dextrogyrate, and from 162.5° to 190° C (324.5° to 374° F) fractions (amounting to 8.52 p.c.) which increased in laevotation with the boiling point, namely from −0.8° to −10.3°. Neither the *BP*, the *USP* nor the *PG* refers to the optical rotation. It is officially stated to be soluble in its own volume of Glacial Acetic Acid. This test has been shown (*PJ* '02, 1 503) by the author and C. M. Carnes to be practically of no value as a test for Oil of Turpentine, although useful as a test of the strength of Glacial Acetic Acid. An acid conforming strictly to the *BP* titration test (which requires a definite figure) cannot be expected to form a clear solution with all samples of Oil of Turpentine when mixed in equal volumes. Commercial samples of Glacial Acetic Acid which require more than the *BP* figure will mix readily without subsequent separation, and most of the commercial acids give a higher figure than the *BP*. With such samples of Oil of Turpentine as had up to that time been examined the mixture of any of them in equal volumes with Glacial Acetic Acid [temperature 14.4° to 16.7° C (58° to 62° F)] became a delicate test for a strength of 99.5 p.c. acid or stronger. The test is also referred to under *Acidum Aceticum Glaciale*.

The more generally occurring impurities are Petroleum, Paraffin Oils, Rosin, Rosin Oil, Petroleum Benzin, Keosene Oil or similar hydrocarbons. Petroleum, Paraffin Oils or Rosin, if present, may be detected by the residue test. Keosene or Rosin Oil, if present, by the evaporation test. Petroleum Benzin, Keosene and similar hydrocarbons by the Sulphuric Acid test, each of which tests is described in small type below. Some work done in the laboratory of the Canadian Inland Revenue Department (*CD* '02, 1 955) has resulted in the following definition of Oil of Turpentine, which must, however, be regarded as provisional, and subject to correction and amplification, it should be colourless, in thin layers, clear, but made decidedly opaque by shaking with 1.0 p.c. of Water and giving an opaque distillate of one-tenth volume which settles clear in a few hours. The peculiar and characteristic odour quite distinct from that of Gasolene, Rosin Oil, or Acetone. It has a sp. gr. between 0.860 and 0.880 (usually about 0.870). Samples which have been long exposed to the air

have a higher density. The first 10 p c fraction has a sp gr of between 0.856 and 0.870 (usually about 0.860), the residual tenth should not exceed 0.900. The boiling point should be between 154° and 158° C (309.2° and 316.4° F), nine-tenths should distil below 180° C (356° F). Fixed residue should not exceed 2 p c, flash point about 32° C (89.6° F). The optical activity of the first fraction should increase in a plus direction by oxidation. The refractive index at 20° C should be between 1.4667 and 1.4722, that of the first fraction should not exceed 1.470. Moistened Starch Iodide paper should not become blue when suspended over Turpentine exposed to the air, free Bromine in solution should not be decolorised. Strong Sulphuric Acid should polymerise and char the sample at a boiling temperature, a rise of temperature should result on mixing with Sulphuric Acid.

Residue—After distillation it should leave little or no residue, *B P*, 1 c c evaporated in a small dish on a water bath should leave not more than a very slight residue, *U S P*.

Evaporation Test—3 drops of Oil of Turpentine placed on a sheet of clean white filter paper and exposed to the air should evaporate entirely without leaving a permanent stain, *U S P*.

Potassium Hydroxide—If 5 c c of the Oil be shaken with an equal volume of Potassium Hydroxide T S, its colour should not become darker than a light straw yellow upon standing 24 hours, *U S P*.

Sulphuric Acid—If 5 c c of the Oil be placed in a small beaker and 20 c c of Sulphuric Acid be gradually added, with agitation, while the beaker is cooled by immersion in cold Water, and the contents, after cooling and renewed agitation, be transferred to a burette, graduated in tenths, the clear layer which forms after the dark mass has settled should not measure more than 0.35 c c (absence of Petroleum Benzine, Kerosene, or similar hydrocarbons), *U S P*.

Preparations

LINIMENTUM TEREBINTHINÆ LINIMENT OF TURPENTINE

Dissolve 1 of Camphor in 13 of Oil of Turpentine and add them gradually to a mixture of 1½ of Soft Soap in 2 of Distilled Water, with constant trituration until a cream is produced, and add Distilled Water, *q s* to yield 20 (about 1 in 1½)

Official in U S, Resin Cerate 55, Oil of Turpentine 35

LINIMENTUM TEREBINTHINÆ ACETICUM. LINIMENT OF TURPENTINE AND ACETIC ACID

Oil of Turpentine, 4, Glacial Acetic Acid (by weight), 1, Liniment of Camphor, 4 (about 1 in 2)

An imitation of St. John Long's celebrated Liniment

Foreign Pharmacopœias—Official in Swed (*Linimentum Terebinthinæ Acetatum*), 9 Oil in 20, Swiss (*Linimentum Terebinthinæ Compositum*), about 3 Oil in 10

Not Official

CONFECTIO TEREBINTHINÆ—Oil of Turpentine, 1 fl oz, Liquorice Root, in powder, 1 oz, Clarified Honey, 2 oz. Rub the Oil of Turpentine with the Liquorice, add the Honey, and mix to a uniform consistence—*B P* 1885

This has been incorporated in the *B P C*

EMULSUM OLEI TEREBINTHINÆ —Rectified Oil of Turpentine, 1b; Expressed Oil of Almond, 5, Syrup, 25, Acacia, 15, Water, *q s* to make 100 — *U S P*

ENEMA TEREBINTHINÆ —Oil of Turpentine, 1 fl oz, Mucilage of Starch, 15 fl oz — *B P* 1885

Oil of Turpentine, $\frac{1}{2}$ to $\frac{1}{2}$ fl oz, Mucilage of Starch, $\frac{1}{2}$ to 1 pint — *St Thomas's*

Oil of Turpentine 2, Mucilage of Starch 100 — *B P C*

LINIMENTUM TEREBINTHINÆ —*Rosin Cerate 65, Oil of Turpentine, by weight, 35 Dissolve the melted Cerate in the Oil of Turpentine and mix thoroughly — *U S P*

*Ceratum Resinæ —Rosin 35, Yellow Wax 15, Lard 50 — *U S P*

UNGUENTUM TEREBINTHINÆ —Oil of Turpentine, 1 fl oz, Resin, in coarse powder, 54 grains, Yellow Wax, $\frac{1}{2}$ oz, Prepared Lard, $\frac{1}{2}$ oz — *B P* 1885

This has been incorporated in the *B P C* as follows —

Oil of Turpentine, by weight, 45, Resin, in coarse powder, 5, Yellow Beeswax 25, Lard 25

Turpentine 1, Yellow Wax 1, Oil of Turpentine, by weight, 1 — *Ger*

VASOLIMENTUM and PAROGENUM TEREBINTHINÆ See p 717

Not Official

THALLINÆ SULPHAS



A yellowish-white crystalline powder, with an odour resembling that of Coumarin, and an aromatic bitter taste

The Sulphate of a synthetically prepared base derived from Chinoline, the full name of which is Tetrahydroparaquamanisol or Tetrahydroparamethoxychinolin

The free base is precipitated from solutions by alkalis, and from it are obtained the Iodide and other Iodinated compounds (*e g*, Periodotetrahydroparamethoxychinolinum) which have been used in cancer

Solubility — 1 in 7 of Water

Medicinal Properties —Antipyretic and antiseptic Has been recommended internally in typhoid and other fevers — *L* '84, 11 1018, *L M R* '85, 456, *B M J* '87, 11 1438

For gonorrhœa, an injection $2\frac{1}{2}$ grains in 150 minims of Water, a bougie 2 grains in 40 grains of Cacao Butter — *B M J* '87, 11 1438, *L M R* '87, 162

Adverse results in gonorrhœa — *B M J* '89, 1 1458

Dose — 3 to 8 grains = 0.2 to 0.52 gramme

Tests —Thalline Sulphate dissolves readily in Water, forming a solution which possesses an acid reaction towards blue Litmus paper, and which becomes brown on exposure to the light From this solution Iodine Solution throws down a brownish-red precipitate, Tannic Acid Solution a white precipitate, and Potassio-mercuric Iodide (Mayer's) Solution a yellow precipitate The dilute aqueous solution, on the addition of Ferric Chloride T.S. a green coloration (changing to a deep red, this green coloration is destroyed by red. lit. soln. ammonia Solution precipitates the free base as a white precipitate which is soluble in Ether The aqueous solution affords with Barium Chloride Solution a white precipitate, which is soluble in Hydrochloric Acid The salt should dissolve to form a colorless solution in Sulphuric Acid, indicating the absence of organic impurities It should leave no weighable residue when heated with free access of air

Cereoli (Antrophore-) are medicated bougies containing a spiral spring wound with fine wire, and coated first with an insoluble layer of White Gelatin,

and then with a diluted Mucilage. They are sometimes medicated throughout and sometimes only medicated externally.

No special medicament is specified in the *Ph Ger*, but they may be medicated in any desired manner.

Antiphores of the salt, described above, have been found useful in gonorrhea.

Foreign Pharmacopœias—Official in Ger

THEOBROMATIS OLEUM

OIL OF THEOBROMA

B P Syn—CACAO BUTTER

A pale yellow or whitish yellow fatty solid, having a distinctive odour of Cocoa and a bland agreeable taste. It is officially described as a concrete Oil, obtained by pressing the warm crushed seeds of *Theobroma Cacao*, L., the *U S P* describes it as a fixed Oil expressed from the roasted seeds of *Theobroma Cacao*. The *P G* describes it as the expressed fat from the seeds of *Theobroma Cacao* free from husk.

Official Preparations—Contained in all the suppositories except Glycerin.

Not Official—Theobromina, Theobrominae Salicylas, Uroclitral, Diuretin and Theocin, Theocin Sodium Acetate, Theobroma Solution Tablet Excipient, Ether-Alcohol Solution of Theobroma Excipient for Tablets.

Foreign Pharmacopœias—Official in Austri, Belg, Dan, Dutch, Ger, Hung, Jap, Norw, Russ, Swed and Swiss (*Oleum Cacao*), Fr (*Beurre de Cacao*), Ital (*Burro di Cacao*), Mex (*Manteca de Cacao*), Port (*Oleo de Cacao*), Span (*Acetate de Cacao*), U S (*Oleum Theobromæ*).

It has been shown (*C D '89*, 1 800) that a large number of substances used in the form of suppositories caused the m p of the mixture to be several degrees higher than the base employed.

Cocoa nut Stearin is sometimes a better substance than Cacao Butter for making suppositories. See p 1154.

Tests—Theobroma Oil softens at 30° to 34° C (86° to 93 2° F), and melts between 31 1° and 33 3° C (88° and 92° F), the *U S P* gives the m p as 30° to 35° C (86° to 95° F), the *B P* gives 31 1° to 33 9° C (88° to 93° F). The m p has been shown to depend largely upon the method by which it is taken, the degree of heat to which the Oil is subjected previous to the determination, the diameter of bore of the capillary tube, and the time allowed to elapse between the melting of the Oil and the actual determination of its m p. It requires about 24 hours in a capillary tube to regain its original m p. Neither the *B P* nor the *P G* refers to the sp gr. The *U S P* gives the sp gr as 0 970 to 0 976 at 25° C (77° F). It usually possesses a sp gr of about 0 990, but authorities differ greatly respecting this constant. It possesses an Acid value of nil to 4 1. A Saponification value of 188 to 198. Neither the *B P*, the *U S P*, nor the *P G* gives a figure for the Acid value. The *U S P* gives 188 to 195 for the Saponification value, neither the *B P* nor the *P G* refers to this latter test. The Iodine value varies from 33 to 37, it is not referred to by the *B P*. The *U S P* states not less

than 33 nor more than 38, the *P G* not less than 34 and not more than 38. 12 samples examined in the author's laboratory possessed Acid values ranging from nil to 4.1, with an average of 2.3, Saponification values ranging from 193.7 to 202.2, with an average of 197.2, and Iodine values ranging from 30.5 to 40.6, with an average of 36.1.

The more generally occurring impurities are Wax, Stearin, Tallow or Suet, or fixed Oils, *e.g.*, Sesame. These may be detected by their influence on the physical constants of the Oil and by their effect on the Acid, Saponification, and Iodine values. Wax, Stearin, Tallow or Suet and other fats may also be detected by the Ether test described in the small type below. The addition of Paraffin Wax will reduce the Saponification value.

Ether—Dissolve 1 gramme of the Oil in 3 c.c. of Ether in a test-tube at 17° C (62.6° F) and place the tube (plunge it frequently, *U S P*) in Water at 82° F (30° C). The liquid should not become turbid nor deposit a granular mass (white flakes, *U S P*) in less than 3 minutes, and if the mixture after congealing be exposed to a temperature of 60° F (15.5° C), it should gradually become clear (absence of Wax, Stearin, Tallow, etc.), *B P* and *U S P* the temperature given in the latter Pharmacopœia is 15° C (59° F), a solution of the Oil in 2 parts of Ether should not become turbid in the course of a day at 12° to 15° C (53.6° to 59° F), *P G*.

Saponification—When saponified by Alcoholic Potassium Hydroxide T.S. it should show a Saponification value of 188 to 195, *U S P*.

Iodine Absorption—If 0.8 gramme of the Oil be dissolved in 10 c.c. of Chloroform in a 250 c.c. bottle or flask, and 25 c.c. of a mixture of equal volumes of Alcoholic Iodine T.S. and Alcoholic Mercuric Chloride T.S. added, and if, after standing for 4 hours protected from light, 20 c.c. of Potassium Iodide T.S. be added and the mixture diluted with 50 c.c. of Water, on titrating the excess of Iodine with Tenth-normal Volumetric Sodium Thiosulphate Solution an Iodine value of not less than 33 nor more than 38 should be obtained, *U S P*, when 1 gramme of the Oil is dissolved in 15 c.c. of Chloroform and mixed with 25 c.c. Alcoholic Iodine Solution and Alcoholic Mercuric Chloride Solution and allowed to remain at rest protected from direct daylight for 4 hours and a solution of 1.5 grammes of Potassium Iodide in 100 c.c. of Water is then added, the mixture when titrated with Tenth-normal Volumetric Sodium Thiosulphate Solution shall show an absorption value of not less than 34 nor more than 38, *P G*.

Not Official

THEOBROMINA Dimethyl-xanthine, $C_7H_8N_2O$, eq. 178.89—White crystalline powder, appearing under the microscope as trimetric needles.

Solubility—1 in 1700 of Water, 1 in 5000 of Alcohol (90 p.c.)

It is the alkaloid contained in the Cacao seeds and is isomeric with Theophylline and Paraxanthine. It is the lower homologue of Caffeine, and has a similar physiological action but stronger. It is much less soluble in Water than Caffeine, and acts the part of a weak Acid, forming compounds with alkalis. The seeds contain 1 to 2 p.c. of the alkaloid.

Diuretic, acting most efficiently in cases of cardiac disease.—*T' G* '93, 767, *B M J E* '93, ii 104. Considered in many respects superior to Diuretin.—*P*, li 299. Diuresis may be prolonged by the subsequent administration of Digitalin ($\frac{1}{12}$ and $\frac{1}{4}$ grain).—*T' G* '96, 330, *L* '96, i 205, ii 1820, *P J* '95, ii 391.

Not a genuine diuretic, but a cardiac stimulant, useful in arterio-sclerosis and aortic incompetence, but the action is temporary and palliative rather than curative.—*B M J E* '05, i 4.

Dose—5 to 10 grains = 0.32 to 0.65 gramme.

Official in Austr, Dutch, Fr, Span, Swed and Swiss

Tests—Theobromine sublimes without decomposition or previous fusion at 290°C (554°F). *Ph Codex* (1908) gives about 260°C (500°F). It dissolves very sparingly in Water. It dissolves in acids and is reprecipitated from solution by alkali, but is soluble in excess of Ammonia Solution or in solutions of Potassium or Sodium Hydroxide. An aqueous solution of Theobromine affords with Mercuric Chloride Solution a white crystalline precipitate. On the addition of Silver Nitrate Solution to a dilute aqueous solution of Theobromine or a Theobromine salt, silver white needles are precipitated after a short time. When a small quantity of Theobromine is evaporated to dryness on a water bath with an excess of Chlorine Water it leaves a reddish brown residue, which assumes a purple violet coloration when moistened with Ammonia Solution. Theobromine may be completely extracted from its solutions by shaking with Chloroform. When ignited with free access of air it should leave no weighable residue.

Theobrominæ Salicylas Theobromine Salicylate may be prepared by dissolving molecular proportions of Theobromine and Salicylic Acid in Water, evaporating to dryness and powdering the residue. The salt is stated to be more stable than the double salt, Sodium Theobromine Salicylate which is decomposed even by Carbon Dioxide. Whilst admitting the advantage which it possesses with regard to its stability, an insurmountable obstacle is presented to its extended use on account of its insolubility.

Tests—Theobromine Salicylate is only very slightly soluble in Water. The solution affords a violet coloration on the addition of Ferric Chloride TS. The salt dissolves in Sodium Hydroxide Solution with the formation of a double Salicylate (Sodium Theobromine Salicylate).

UROCITRAL (Theobromine Sodium Citrate)—A white powder, soluble in warm Water. It is stated to contain 45 p.c. Theobromine. Introduced as a diuretic.—*B M J* '05, 1, 81.

DIURETIN Sodium Theobromine Salicylate $\text{C}_7\text{H}_5\text{N}_3\text{O}_7, \text{C}_6\text{H}_5\text{O}_2, \text{Na}$, eq 359.66—A white, odourless, unstable powder.

It should be kept in well stoppered glass bottles of a dark amber tint and exposed as little as possible to contact with the air, as it is liable to absorb Carbon Dioxide, decomposition simultaneously occurring.

A comparison of this drug with Agurin shows (*B M J E* '04, 11, 59) little fundamental difference of action, but only difference of degree.

Dose—10 to 20 grains = 0.65 to 1.3 grammes, thrice daily.

Ph (Gr) maximum single dose, 1 gramme maximum daily dose, 6 grammes.

Foreign Pharmacopœias—Official in Austr, Belg, Dan, Dutch, Ger, Ital, Jap, Mex, Span, Swed and Swiss.

Tests—Sodium Theobromine Salicylate dissolves, when freshly prepared readily in Water, forming a colourless solution possessing an alkaline reaction towards red Litmus paper. A diluted aqueous solution when acidified with Acetic Acid yields on the addition of Ferric Chloride TS a violet coloration. When the aqueous solution is acidified with Hydrochloric Acid, Salicylic Acid as well as Theobromine is precipitated as a white precipitate, redissolving on the addition of Sodium Hydroxide Solution, but not on the addition of Ammonia Solution. 10 c.c. of a 1 in 5 w/v aqueous solution, from which the Theobromine and the Salicylic Acid have been precipitated by the addition of Hydrochloric Acid, and again redissolved by the addition of Sodium Hydroxide Solution (15 p.c.), when shaken with 10 c.c. of Chloroform and the Chloroform evaporated to dryness shall leave not more than 0.005 gramme of residue for each 1 gramme of Sodium Theobromine Salicylate represented in the original volume employed. A weighed quantity of 2 grammes of Sodium Theobromine Salicylate is dissolved by the aid of a gentle heat in a porcelain dish in 10 c.c. of Water the solution is mixed with 5 c.c. of a sufficient quantity of Normal Volumetric Hydrochloric Acid Solution to render it slightly acid, and when mixed, 1 drop of a diluted Ammonia Solution (1 to 10) is added, and the very faintly alkaline mixture is allowed to stand for 3 hours at a temperature of from 15° to 20°C (59° to 68°F).

with intervals of frequent stirring, the test is filtered through a tared filter paper, previously dried at 100°C , and dried twice with 10 c c of cold Water, dried till constant at 100°C (212°F) and when cooled weighed. The weight shall amount to at least 0.8 grammes, corresponding to at least 40 p c of Theobromine. 1 part by weight of this precipitate mixed with 100 parts of Chlorine Water and evaporated to dryness on a water-bath leaves a yellowish-red residue, which, on the addition of a little Ammonia Water, yields a beautiful purple coloration. The filtrate from this Theobromine residue is acidified with Salicylic Acid, which may be determined by acidulating with Salicylic Acid and shaking out with Chloroform. The chloroformic solution is washed with Water till free from mineral acid, sufficient Water added to form a separate layer, a few drops of Phenolphthalein Solution added and the mixture titrated with Tenth-normal Volumetric Sodium Hydroxide Solution, 1 c c of Tenth-normal Volumetric Sodium Hydroxide Solution corresponds to 0.013701 grammes of Salicylic Acid, it should contain about 38.5 p c.

THEOCIN Theophylline, Dimethylxanthin $\text{C}_7\text{H}_8\text{N}_2\text{O}_2$, eq. 178.89—Colourless, or white crystalline needles possessing a bitter taste. It is isomeric with Theobromine and Paraxanthin. It is stated to crystallise with 1 molecule of Water of crystallisation, which it loses at 110°C (230°F). Soluble 1 in 190 Water, 1 in 80 of Alcohol (90 p c), forming Potassium and Ammonium compounds which are readily soluble. It is a synthetic alkaloid, and is identical in composition with Theophylline, the alkaloid occurring with Theine or Caffeine in tea. It has been introduced as a diuretic. It has been used in kidney disorders with general dropsy, and appears to be most efficient when considerable cedema exists—*B M J E* '03, ii 39, 56, *A J P* '03, 27, *C D* '03, i 50, *P J* '03, i 2. In *B M J* '05, i 1079 it is pointed out that it is a less powerful stimulant than Caffeine, but more active as a diuretic than Caffeine or Theobromine. A full dose may cause nausea or even vomiting, therefore the dose should not exceed 6 or 7 grains = 0.4 to 0.6 grammes.

Dose—3 to 6 grains = 0.2 to 0.4 grammes.

It is also supplied in tablet form, each tablet containing 4 grains = 0.26 grammes.

Tests—Theophylline melts at about 264°C (507°F). Synthetic Theocin melts at 268°C (514°F). It dissolves readily in warm Water, but is only sparingly soluble in cold Alcohol (90 p c), it is readily soluble in very dilute Ammonia Solution. When evaporated to dryness with Chlorine Water it yields a scarlet residue, changing to purple-red on the addition of a little Ammonia Solution. When ignited with free access of air it should leave no weighable residue. On the addition of Silver Nitrate to an aqueous solution of Theophylline an amorphous precipitate is produced.

THEOCIN SODIUM ACETATE $\text{C}_7\text{H}_7\text{NaN}_2\text{O}_2 \cdot \text{NaC}_2\text{H}_3\text{O}_2$, eq. 282.23—A white powder containing about 65 p c of anhydrous Theocin. It is soluble 1 in 6 of Water, 1 in 80 of Alcohol (90 p c), and insoluble in Ether. It is a double salt of Sodium Acetate, and 1.3 Dimethylxanthin Sodium. It was introduced as a diuretic and is indicated in all forms of dropsy in which the functions of the kidneys are not too seriously impaired by disease. It may be administered in cases of cedema resulting from renal disease, with the exception of cases of glomerulo-nephritis, in which active interference is contra-indicated. It should afford beneficial results in interstitial nephritis and in arterio-sclerosis of the kidneys and in granular contracted kidney, and has been used with benefit for its vaso-dilator and antispasmodic effects in angina pectoris.

Striking effect as a diuretic, but to be effective must be given with a cardiac tonic such as Digitalis, in 3 to 8-grain cachets every 4 hours, effects to be carefully watched, as it is apt to irritate the stomach—*B M J* '07, ii 388.

Confirmation of the foregoing, in a severe case of ascites and cedema, no ill-effects—*B M J* '07, ii 752.

Dose—4 grains = 0.26 of a grammes.

Tests—Theocin Sodium Acetate dissolves readily in Water, forming a solution which is slightly alkaline in reaction towards blue Litmus paper. It affords with Ferric Chloride T.S. a dark red coloration, and on boiling a brownish-

red precipitate. When heated with Sulphuric Acid it evolves a strong characteristic acetous odour, when further warmed with a little Alcohol (90 p c) it evolves a characteristic odour of Ethyl Acetate (Acetic Ether). When ignited with free access of air it leaves a residue which when dissolved in Water possesses a strong alkaline reaction towards red Litmus paper, and effervesces on the addition of diluted Hydrochloric Acid, it yields the tests distinctive of Sodium given under that heading.

THEOBROMA SOLUTION TABLET EXCIPIENT—Oil of Theobroma, 25, Hard Soap, 5, Powdered Tragacanth, 0.5, Benzoic Acid, 0.25, Water, to 100. Dissolve the Soap in 25 parts of Water by heat, add the hot solution to the melted Theobroma and mix by whisking or agitation, shake in the Tragacanth, add the rest of the Benzoic Acid, then the rest of the Water. Gum Acacia may be used in place of Soap without making any appreciable difference in the general utility of the product, in either case it should be a thick, smooth, white cream, free from lumps.—*C D '03, 11 231*

This has been incorporated in the *B P C* under the title *Emulsio Theobromatis*, using 3 of Hard Soap in the place of 5 given originally.

ETHER-ALCOHOL SOLUTION OF THEOBROMA EXCIPIENT FOR TABLETS—Oil of Theobroma, 1, Ether, to 6 fl oz.—Dissolve, and add an equal volume of Rectified Spirit as required for use.—*C D '03, 11 231*

This has been incorporated in the *B P C* under the title *Liquor Theobromatis Aethereus*.

For general directions for making Compressed Tablets see p. 1190

THUS AMERICANUM.

FRANKINCENSE

A softish, pale, opaque solid, possessing an agreeable terebinthinate odour. On keeping it hardens and forms a translucent brittle solid. It is officially described as the concrete oleo resin which is scraped off the trunks of *Pinus palustris*, Mill, and *Pinus Teda*, L.

From the Southern States of North America

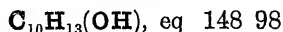
Solubility—Almost wholly soluble 1 in 1 of Alcohol (90 p c), entirely 4 in 3 of Ether.

Medicinal Properties—Used externally for the same purposes as Resin.

Official Preparation—Used in the preparation of *Emplastrum Picis*.

THYMOL.

THYMOL



Large, colourless, translucent, oblique, rhombic prisms, having a distinctive somewhat agreeable, Thyme-like odour, and burning, aromatic taste. It is a crystalline Phenol contained in the volatile Oils of *Thymus vulgaris*, L, *Monarda punctata*, L, and *Carum Coptivum*, Benth and Hook f, but is chiefly obtained commercially from the last named. Thymol is described by the *USP* as a Phenol occurring in the volatile Oil of *Thymus vulgaris*, L, and in some other volatile Oils.

It should be kept in well-closed vessels of a dark amber tint.

Solubility.—1 in 1500 of Water, 1 in 190 of Glycerin, 8 in 3 of Alcohol (90 p.c.) or Ether, 8 in 5 of Chloroform, 1 in 6 of Petroleum Spirit, 1 in 3 of Oil of Turpentine, 1 in 2 of Olive Oil, 4 in 3 of Glacial Acetic Acid, 1 in 6 of Solution of Potassium Hydroxide

The above figures for solubility have been incorporated in the *B.P.C.* The expressions peculiar to the *Companion*, 8 in 3 of Alcohol (90 p.c.) or Ether, 8 in 5 of Chloroform, and 4 in 3 of Glacial Acetic Acid being also used

Medicinal Properties—A saturated solution in Water is a very powerful antiseptic, used as an intestinal antiseptic in diarrhoea and typhoid. As an ointment or soap in parasitic skin diseases. As an inhalation in laryngitis and bronchial affections, and for many other conditions in which Carbolic Acid is useful. It is a very powerful deodorant, and is a local anæsthetic.

In ankylostomiasis, no vermifuge is comparable to it. 10 to 60 grains for fairly robust patients, not more than 10 grains for those who are very ill, or much advanced in years.—*B.M.J.* '03, 1 720

Recommended in ankylostomiasis (*P.* lxviii 685), in 2 or 3 doses of 2 grammes = 30 grains, at 2 hours' interval, after a little Coffee or Broth in the early morning, and after a Calomel and Senna purge. 6 grammes is the limit advised, and a dose of Epsom salts should be given 2 hours after the last dose to eliminate the Thymol from the intestinal tract. Children, $\frac{1}{4}$ to $\frac{1}{2}$ the full dose. The administration of any of the usual solvents of Thymol must be avoided.

In ankylostomiasis should be given (*L.* '05, 1 860) in large and repeated doses, $\frac{1}{2}$ drm. every 2 hours for several doses.

Usually employed as a deodorant, which property it possesses to a marked degree, its aqueous solution is very useful in a night commode, and an extremely small quantity of it will keep urine, when it is required to make a 24 hours' collection for analytical purposes.

Dose— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Not Official—Glycerinum Thymol Alkalinum, Glycerinum Thymol Compositum, Liquor Antisepticus, Liquor Thymol, Thymol Antiseptic Dressing, Unguentum Thymol, Vapor Thymol, Oleum Thymi, Aristol, Carvacrol Iodide and Thymol Carbonate

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Jap., Mex., Norw., Russ., Span., Swed., Swiss and U.S. Span. and U.S. have also Thymolis Iodidum

Tests.—Thymol melts at about 50° C (122° F), the *U.S.P.* and the *P.G.* give 50° to 51° C (122° to 123.8° F), the *B.P.* does not include a m.p. The sp. gr. is given by the *U.S.P.* as 1.030 at 25° C (77° F). The *U.S.P.* also states that when liquefied by fusion it is lighter than Water. The *B.P.* states that the crystals sink in cold Water, but at a temperature of 43.3° to 51.7° C (110° to 125° F) the crystals melt and rise to the surface. The *P.G.* states that the crystals sink in Water, but that melted Thymol floats on the surface of Water. Neither the *B.P.* nor the *U.S.P.* refers to the boiling point. It boils at 232° C (449.6° F). The *P.G.* states that it boils at 228° to 230° C (442.4° to 446° F). When mixed with an equal proportion of Camphor, Menthol or Chloral Hydrate it liquefies. It dissolves only sparingly in Water, but readily in Alcohol (90 p.c.), the alcoholic solution being optically inactive. The *B.P.* states that a solution of Thymol in half its bulk of Glacial Acetic Acid warmed with an equal volume of Sulphuric Acid yields a reddish-

violet colour. The *USP* and the *PG* state that a very small crystal of Thymol dissolved in 1 cc of Glacial Acetic Acid yields a liquid which on the addition of 6 drops of Sulphuric Acid and 1 drop of Nitric Acid yields a deep bluish green colour when viewed by reflected light. It dissolves in 4 parts by weight of Sulphuric Acid at the ordinary temperature, forming a yellowish liquid which on gentle warming assumes a beautiful rose-red coloration. If this solution be poured into 10 volumes of Water, and the mixture allowed to stand with frequent intervals of shaking at a temperature of 35° to 40° C (95° to 104° F) in contact with an excess of Lead Carbonate, and the liquid be then filtered it yields a filtrate which on the addition of a little Ferric Chloride Solution assumes a beautiful violet colour. A distinctive reaction for Thymol and also for Carvacrol is the reddish violet coloration produced on heating a small quantity of either with 0.1 of a gramme of Potassium Hydroxide and 20 drops of Chloroform. Thymol requires the addition of a few drops of Alcohol to effect solution before the colour is produced. The *USP* describes a somewhat similar test, 1 gramme of Thymol heated in a test-tube in a water bath with 5 cc of Sodium Hydroxide Solution (10 p.c.) affords a clear colourless solution possessing a very slight reddish tint, becoming darker on standing, but without the separation of oily drops. On the addition of a few drops of Chloroform to this liquid it affords when agitated a violet coloration.

The more generally occurring impurities are Phenol, non volatile organic impurities, and inorganic impurities. Phenol may be detected by the Ferric Chloride and the Bromine tests described in small type below. Non volatile organic impurities and inorganic impurities by the residue test also described below.

Residue—It is completely volatilised at the temperature of a water bath, *BP* and *USP*, 0.1 gramme volatilised on a water bath should not leave a weighable residue, *PG*.

Ferric Chloride—An alcoholic solution of Thymol should not be coloured by T.S. of Ferric Chloride, *USP*, an aqueous solution should be neutral and should not be coloured violet, *PG*.

Bromine—In an aqueous solution of Thymol, Bromine Water should produce a milky turbidity but not a crystalline precipitate, *PG*.

Not Official

GLYCERINUM THYMOL ALKALINUM—Sodium Bicarbonate, 100 grains, Sodium Borate 200 grains, Sodium Benzoate, 80 grains, Sodium Salicylate, 40 grains, Menthol, 2 grains, Pumpkin Seed Oil, 4 minims, Wintergreen Oil, 2 minims, Thymol, 4 grains, Eucalyptol, 12 minims, Alcohol (90 p.c.) 4 fl. oz., Glycerin, 2 fl. oz. Solution of Camphor, 40 minims, Distilled Water, *qs* to produce 20 fl. oz. Dissolve the salts in the Water, add the Glycerin and Solution of Camphor, then add the Oils previously dissolved in the Alcohol, and filter—*Bourne's Formula*.

This has been incorporated in the *BPC* under the title **Glycerinum Thymol Compositum**. *Syn* Glycerinum Thymol Alkalinum, as follows—Sodium Bicarbonate, 1, Sodium Borate, 2, Sodium Benzoate, 0.75, Sodium Salicylate, 0.50, Menthol, 0.03, Oil of Pine, 0.05, Oil of Wintergreen, 0.03, Thymol, 0.05, Eucalyptol, 0.13, Alcohol (90 p.c.), 2.50, Glycerin, 10, Solution of Camphor, 0.50, Distilled Water, *qs* to produce 100,

LIQUOR ANTISEPTICUS—Boric Acid, 2, Benzoic Acid, 0.1, Thymol, 0.1, Eucalyptol, 0.025, Oil of Peppermint, 0.05, Oil of Gaultheria, 0.025, Oil of Thyme, 0.01, Alcohol (95 p c), 25, Purified Talc, 2, Water, *q s* to make 100. Dissolve the Boric Acid in 70 of Water and the Benzoic Acid in 15 of Alcohol, pour the aqueous solution into the alcoholic solution. Then dissolve in a mortar the Thymol in the Oils, incorporate thoroughly the Purified Talc, and add with constant trituration the solution first prepared. Allow the mixture to stand for 48 hours with occasional agitation, filter, add 10 of Alcohol to the clear filtrate, and sufficient Water to make 100—*U S P*

This has been incorporated in the *B P C* under the title of **Liquor Thymolis Compositus** with *synonym* Liquor Antisepticus, using 26.50 of Alcohol (90 p c) in place of 25 of Alcohol (95 p c)

Liquor Antisepticus (Volckmann)—Thymol, 1, Alcohol (90 p c), 10, Glycerin, 20, Distilled Water, 100

LIQUOR THYMOL—Thymol, 1, Alcohol (90 p c), 100. This solution is very useful, as it may be diluted to any extent with Water without precipitation. Half a pint diluted to a gallon is about the same strength as a saturated aqueous solution.

THYMOL ANTISEPTIC DRESSINGS—Gauze, 5 p c, and Wool, 5 p c

UNGUENTUM THYMOL—Thymol, 20 grains, Soft Paraffin, 1 oz—*London*

VAPOR THYMOL—Thymol, 6 grains, Alcohol (90 p c), 60 minims, Light Magnesium Carbonate, 3 grains, Water, to 1 fl oz—*Throat*

A teaspoonful in a pint of Water at 140° F for each **inhalation**

OLEUM THYMI—The rectified Oil forms an almost colourless or yellow oily liquid, having a pleasant aromatic Thyme-like odour and a sharp aromatic taste. The crude Oil is a reddish or reddish-brown oily liquid possessing similar characteristics of taste and odour. It is the Oil distilled, principally from the fresh flowering herb, *Thymus vulgaris*. Should contain from 25 to 35 p c of Phenols (Thymol and Carvacrol). The rectified Oil soon darkens in colour on exposure to air and light, and should be kept in well-stoppered bottles of a dark amber tint.

The Oil is not official in the *B P*. The *U S P* and the *P G* describe it as the volatile Oil distilled from the leaves and flowering tops of *Thymus vulgaris*, both Pharmacopœias require it to contain not less than 20 p c by volume of Phenols.

Foreign Pharmacopœias—Official in Fr, Ger, Jap, Russ, Span, Swiss and U S

Tests—Oil of Thyme has a sp gr of 0.900 to 0.930, the *U S P* states 0.900 to 0.930 at 25° C (77° F), the *P G* 0.900. It is slightly laevogyrate, the optical rotation being from -1° to -3°. The *U S P* states not more than -3° in a 100 mm tube at a temperature of 25° C (77° F). The *P G* does not give the optical rotation. It dissolves in half its volume of Alcohol (94.9 p c) and in 1 to 2 volumes of Alcohol (80 p c). The *P G* states that it is soluble in 3 parts by weight of a mixture of 100 parts by volume of Alcohol (90 p c) and 14 parts by volume of Water. The alcoholic solution yields with a drop of Ferric Chloride T.S. a greenish-brown coloration, changing to reddish. It is required by the *U S P* to contain not less than 20 p c by volume of Phenol, determined by measuring the volume of unabsorbed non-volatile Oil remaining after treating the Oil with a 1 in 20 Sodium Hydroxide. A measured quantity of 10 c.c. of the Oil introduced into a burette having a capacity of 50 c.c. and containing 40 c.c. of a 1 in 20 Sodium Hydroxide Solution, the burette is well corked and the mixture shaken thoroughly, and then set aside for from 12 to 24 hours, the drops of Oil adherent to the side of the burette are detached by gentle tapping and rotation. When the alkaline liquid has become clear, the volume of unabsorbed Oil is measured and subtracted from the original amount of Oil taken, the difference multiplied by 10 indicates the percentage of Phenols in the Oil, the unabsorbed Oil should not measure more than 8 c.c.

The *P G* adopts a corresponding limit, but shakes a measured quantity of 5 c c of the Oil with 30 c c of a mixture of 10 c c of Sodium Hydroxide Solution (15 p c w/w), and 20 c c of Water in a graduated cylinder, and allows the mixture to stand until the alkaline solution has become clear, the volume of unabsorbed Oil is then read off. It should be subtracted from the volume of Oil used for the determination (5 c c), and the result multiplied by 20 which indicates the percentage by volume of Phenols present in the Oil. The Oil when shaken with 10 times its volume of hot Water, cooled, and the liquid filtered through a wet filter yields a filtrate which is not coloured bluish or violet by Ferric Chloride T S, indicating the absence of Phenol.

A comparison of commercial Thyme and Origanum Oils is given (*P J '03*, 1 803), French Thyme Oil, from *Thymus vulgaris*, had a sp gr of 0 905 to 0 920 and contained from 18 to 45 p c of Phenol, Wild Thyme Oil, from *Thymus serpyllum* had a sp gr of from 0 890 to 0 905, contained practically no Phenols, a Spanish Thyme Oil, of doubtful origin, had a sp gr of from 0 930 to 0 950, and contained 50 to 70 p c of Phenols. Thyme Oil from *Origanum hirtum* had a sp gr of from 0 940 to 0 980 and contained from 60 to 85 p c of Phenols, Smvrna Oil from *Origanum Smyrnaceum* had a sp gr of 0 915 to 0 945, and contained from 25 to 60 p c of Phenols, Cyprus Oil from *Origanum majoranoides* had a sp gr of 0 961 to 0 967, contained 78 to 84 p c of Phenols, whilst a sample of Sicilian Oil had a sp gr of 0 920, and contained 44 p c of Phenols.

ARISTOL (Thymol Iodide, $C_{10}H_{14}O I_2$, eq 545 76) —A bright yellowish or brownish yellow or reddish yellow bulky powder with a slight aromatic odour somewhat resembling Iodoform. It is insoluble in Water and Glycerin, slightly soluble in Alcohol, readily soluble in Ether and Chloroform. It has been introduced as a substitute for Iodoform. Used successfully as a 10 p c Ointment, or by dusting the powder on ulcerating lupus, tinea, and syphilitic ulcers, in psoriasis and eczema a 10 p c solution in flexible collodion, as a pessary in ulceration of vagina or cervix.

It has been used as a dressing for burns.

Tests —Aristol when heated is decomposed. When heated with concentrated Sulphuric Acid it is decomposed with the separation of Iodine. 1 decigramme when shaken with 20 c c of Water and filtered yields a filtrate which should not become more than opalescent on the addition of Nitric Acid and Silver Nitrate Solution. 5 decigrammes shaken with 10 c c of Water and filtered should afford a filtrate which should impart no blue colour to red Litmus paper, indicating the absence of alkalis. 5 decigrammes shaken with 10 c c of Water and the mixture filtered affords a filtrate which should not be coloured blue upon the addition of Starch Solution, indicating the absence of free Iodine. 0 5 of a gramme when ignited with free access of air should leave not more than 0 015 gramme of residue.

CARVACROL IODIDE —A light yellow or reddish brown powder, insoluble in Water and Alcohol, but soluble in Ether and in Chloroform, produced by the action of Iodine and Potassium Iodide on Carvacrol in solution.

As a germicide it is stated to be 5 times more powerful than Iodoform, and being more bulky is better adapted as a dusting powder. The substance which was prepared for many years by the author at the suggestion of Dr Mortimer Granville is of a reddish brown colour, but more recently a substance of a light yellow colour has been used in Germany as a substitute for Iodoform.

Iodocrol, a fancy name applied to the latter product.

THYMOL CARBONATE (Tyratol) —Forms a tasteless white powder. Recommended as a powerful vermifuge —*C D '01*, 11 344.

Arhovin, a product of Diphenylamine and Thymolbenzoic ester, in gonorrhoea (*B M J E '07*, 1 95), an ideal preparation in gonorrhoea, 1 to 2 p c solution in Olive Oil rapidly increased to 3 or 5 p c —*B M J E '06*, 11 87.

THYROIDEUM SICCUM.**DRY THYROID**

A pale buff-coloured to light brown, somewhat hygroscopic, amorphous powder, possessing a peculiar distinctive meat-like odour. The powdered desiccated Thyroid gland is official in the *USP*, but not in the *PG*. It is described as the Thyroid glands of the sheep (*Ovis aries*), Linné, freed from fat, cleaned, dried and powdered. The *BP* requires the healthy gland to be used, and after drying below 40° C (104° F) extracts the fat with Petroleum Ether, but it is not definitely stated in the *USP* monograph that the healthy gland should be employed, of course this would naturally be inferred, the *USP* states freed from fat, but does not indicate a method. 1 part of the desiccated Thyroid gland is stated to represent approximately 5 parts of the fresh gland.

Medicinal Properties—Has been used with success in myxedema and certain forms of insanity, obesity, goitre and cretinism, psoriasis and chronic scaly skin diseases. Thyroid should never be given in exophthalmic goitre.

Preparations—*BMJ* '92, ii 1384, 1459, *L* '93, i 273, 396, in goitre—*L* '95, ii 169, *BMJ* '95, ii 75, '96, i 48, in cancer—*L* '96, ii 106, 162, in cretinism—*L* '96, i 853, 1446, '97, ii 853, '02, i 1565, *BMJ* '01, i 1143, '02, i 1259, in " '94, i 786, '96, ii 1200, *L* '96, ii 41, 470, in psoriasis—*BMJ* " '95, i 697, *L* '95, i 813, *BMJE* '95, ii 85, ichthyosis—*BMJ* '95, i 696, in pityriasis rubra—*BMJ* '95, i 695, in rickets—*BMJE* '02, i 40.

Oophorectomy combined with administration of thyroid has been recommended in inoperable carcinoma of the breast, in small doses, gradually increased to 15 grains daily—*BMJ* '00, ii 1161, '01, ii 1145, 1439, '02, i 508, *L* '01, ii 388, 966, 967, '02, i 888, *TG* '99, 609.

12 p.c. of recoveries in cases of insanity which were not hopeless, but intractable by ordinary methods. It appears to be more efficacious in women than in men, and the best all-round results were connected with the insanity of child bearing—*BMJ* '00, ii 818.

It powerfully affects the metabolism generally of the body cells, raising their tone and improving their vigour—*BMJ* '01, ii 1147.

A useful general *résumé* of our knowledge of the Thyroid Extract, forming portion of the Hunterian Oration on Organo-Therapeutics—*L* '02, i 1091.

In puerperal eclampsia, 5 grains thrice daily for 6 days, followed by 5 grains every 8 hours for 17 days, an interval of 14 days, and then doses of 5 grains daily—*BMJ* '02, i 1214, *L* '02, i 824, ii 459, '03, i 307.

10 grains given in each case on admission, and 5 grains every 4 hours afterwards in puerperal eclampsia—*L* '04, i 1057, *BMJ* '04, i 895.

In psoriasis—Initial dose should not exceed 5 grains once daily, and increment should be gradual and spread over 2 or 3 weeks, and should seldom exceed 15 grains a day, not giving more to patients who are not daily under observation—*BMJ* '03, i 656, *L* '03, i 785.

In glycosuria, 1½ grains in tablet form 3 times a day—*L* '03, ii 187.

In a number of cases of confirmed epilepsy, in which preparations of the Thyroid gland were given over considerable periods, no appreciable result was detected either in the mental condition or in the frequency or severity of the fits—*L* '05, i 710. It has a marked alterative influence in certain chronic affections of the skin. Its effect in cases of psoriasis is so evanescent as to make it of little practical value. It is of use, however, in quickening the healing of small indolent ulcers—*BMJ* '05, i 700.

Dose.—3 to 10 grains = 0.2 to 0.65 gramme

Official Preparation Liquor Thyroides

Not Official —Elixir Thyroides, Liquor Thyroides, Tablets of Thyroid Gland, Iodothyryn and Thyroglandin

Foreign Pharmacopœias —Official in Belg (Thyroides) and U S

Tests —Dried Thyroid is not officially required to answer any definite chemical tests. The *USP* requires that 1 gramme of the desiccated Thyroid gland when mixed with an equal weight of pure Sodium Hydroxide and carefully fused in a silver dish until a white mass remains, Potassium Nitrate being added during the fusing to assist oxidation, yields, when the fused residue is dissolved in a small quantity of Water, a solution which treated with 2 grammes of Sodium Nitrate acidified with concentrated Nitric Acid and shaken with 5 c c of Chloroform imparts to the chloroformic liquid a decided pink to violet coloration. A cold extract of desiccated Thyroid glands treated with 2 grammes of Sodium Nitrate and acidified with strong Nitric Acid should not give the Iodine test on shaking with Chloroform. A preferable method of performing the test is that suggested (*YBP* 1883, 530, *PJ* '98, 11 546), and Chloroform is not found to be a suitable solvent for the Iodine, the sample is never burnt to ash, but always into Charcoal in the presence of a slight excess of Sodium Hydroxide, the risk of loss of Iodine by adding Potassium Nitrate to promote oxidation never being incurred. To liberate the Iodine from the aqueous solution of the charred residue a few drops (1 to 3) of Nitro Sulphuric Acid are used, the Nitro Sulphuric Acid being prepared by treating Starch with Nitric Acid and passing the Nitrous fumes into the Sulphuric Acid (1.043 sp gr) to saturation. Carbon Bisulphide is employed as a solvent for the liberated Iodine, and the tests are performed in large tubes of even bore and compared with standard solution of Potassium Iodide treated in the same manner. It is claimed by this method $\frac{25.0}{100.0}$ part of Iodine is easily detected and measured, and up to $\frac{10.0}{100.0}$ part the estimation is very accurate. When incinerated the *USP* states that desiccated Thyroid glands should yield not more than 6 p c of ash.

Preparation**LIQUOR THYROIDES** THYROID SOLUTION

A liquid prepared from the fresh and healthy Thyroid gland of the sheep

This preparation does not appear to be a success pharmaceutically, as it readily undergoes decomposition. The menstruum is equal parts of Glycerin and Distilled Water, containing about 1 of Phenol in 400 of the total volume.

Glycerin is stated not to dissolve out Thyroidin —*PJ* '98, 11 167, *CD* '98, 11 288. This statement has been contradicted —*PJ* '98, 11 482. But reaffirmed on strong evidence —*PJ* '98, 11 546.

Tests —When evaporated to dryness, the residue moistened with Sodium Hydroxide Solution and fused, the charred residue extracted with Water, the excess of alkali neutralised and the solution mixed with 1 to 3 drops of Nitro-Sulphuric Acid, as described under Thyroidum Siccum, and shaken with a few c c of Carbon Bisulphide a

decided violet coloration should be imparted to the Carbon Bisulphide Solution. No test for the presence of Iodine compounds is given in the *B P*

Not Official

ELIXIR THYROIDEI (*Squue*) —A clear, aromatic, reddish liquid, containing the entire active principles of the Thyroid gland of the sheep. Each fl dram is equal to $1\frac{1}{2}$ grains of dry Thyroid.

Dose —1 to 2 fl dram = 3 G to 7 2 c c

ELIXIR THYROIDEI (*Amowu*) —Prepared with a Glycerin menstruum, 1 fl oz equivalent to 1 entire sheep's Thyroid gland.

Dose —30 to 60 minims = 1 8 to 3 6 c c

LIQUOR THYROIDEI (*Squue*) —A transparent, pale reddish liquid, containing the entire active principles of the gland. Each fl dram is equal to 6 grains of dry Thyroid.

Dose —10 to 60 minims = 0 6 to 3 6 c c

TABLETS OF THYROID GLAND —Each tablet containing the equivalent of $1\frac{1}{2}$, $2\frac{1}{2}$ 5 or 10 grains of the entire substance of the Thyroid gland. **Tablets**, each containing 5 grains, equivalent to 2 grains of the desiccated substance.

IODOTHYRIN (Thyroidin) —An amorphous light brown powder, insoluble in Water, soluble in Alcohol. Dissolved by alkalis and acids, with the addition of an acid. It is an organic compound of Iodine, a principle of the Thyroid gland, free from albuminoids, adjusted with Sugar of Milk to equal in strength the active substance of the fresh gland, and standardised to contain 0·8 p c of Iodine. Usually standardised by dilution with Milk Sugar, to contain a definite percentage of Iodine —*L* '96, 1 592, 666, 941, '97, 11 855, *B M J* '96, 1 722, *B M J E* '96, 11 59, '97, 11 8, *P J* '96, 1 161, 11 215, 388, '97, 1 287.

Tests —Iodothyryn when moistened with Sodium Hydroxide Solution and carefully charred leaves a carbonaceous residue which when dissolved in Water, the alkali neutralised with diluted acid and the solution treated with Nitro-Sulphuric Acid, as described under Thyroideum Siccum, yields when shaken with Carbon Bisulphide Solution a decided violet coloration.

THYROGLANDIN —A light yellowish-brown or brown, somewhat hygroscopic, amorphous powder, which is stated to consist of the entire active constituents of the gland. It contains the Iodoglobulin obtained from the fresh glands by simple treatment with Water, together with the total amount of Iodothyryn obtained by subsequent treatment of the residual glands with 1 p c Soda Solution and exact neutralisation with Hydrochloric Acid —*P J* '98, 11 167, 654, *C D* '98, 11 288, 970, *B M J* '98, 11 79.

Dose —1 to 5 grains = 0 06 to 0 32 grammes

Tests —Thyroglandin when moistened with Sodium Hydroxide Solution and carefully charred leaves a carbonaceous residue which when dissolved in Water, the alkali neutralised with diluted acid and the solution treated with Nitro-Sulphuric Acid, as described under Thyroideum Siccum, yields when shaken with Carbon Bisulphide Solution a decided violet coloration.

Thyrodoctin is stated (*B M J* '07, 1 756) to be the dried blood of animals from which the Thyroid glands have been removed. A reddish-brown powder, put up in capsules containing 5 grains each.

TINCTURÆ.

TINCTURES

Most of the Tinctures of the British Pharmacopœia are directed to be made either by 'maceration' or by 'percolation', the number in each class is nearly equal, but if anything the latter predominate, about a dozen are made by simple solution, or mixing the ingredients

The official directions for maceration and percolation are much the same as in 1864 for percolation the ingredients are macerated with a portion of the menstruum for 48 hours, and then percolated with more of the same, the marc is pressed and the whole yield of liquid made up to the required volume, for maceration, the ingredients are mixed with the required quantity of menstruum, and after 7 days strained, pressed and if necessary the liquid is filtered, in 1864, 1867, and 1885, the macerated tinctures were finally made up to a volume, but in 1898 this was omitted

The degrees of comminution appeared first in the 1885 edition

The following *B P* Tinctures are standardised —Cinchona, Jalap, and Opium the Tinctures of Belladonna and Nux Vomica are made from standardised Fluid Extracts, Ammoniated Tincture of Opium and Compound Tincture of Camphor are made from standardised Tincture of Opium, Compound Tincture of Cinchona from standardised Tincture of Cinchona

The strengths of the various Tinctures have been adjusted so as to have a dosage of 5 to 15 minims for the potent Tinctures, and 30 to 60 for the less potent

With regard to the Tinctures contained in the Continental Pharmacopœias a comparison is given under each separate Tincture in the *Companion* paragraphs commencing *Foreign Pharmacopœias* The Potent Tinctures given therein are compared with the standards adopted by the *Brussels Conference* and the alcoholic strength of the Tincture is also given

The tabulated comparison of the chief standardised potent preparations of the British, United States, German and French Pharmacopœias given at the commencement of this book shows at a glance the alkaloidal strengths and the standards for the Tinctures official in the four Pharmacopœias with which the present volume is chiefly concerned, and which are probably of the most material interest to English readers

The Tinctures or Teintures Alcooliques of the *F. Codez* (1908) are liquid medicaments resulting from the solvent action of Alcohol on various substances, they consist of 'simple' or 'compound' Tinctures, simple being prepared with the single substance, the compound where several substances are used in the preparation They are prepared by maceration or percolation, Alcohol 60 p c, 70 p c, 80 p c, or 95 p c, being employed according to the nature of the drug to be exhausted All simple tinctures of heroic drugs, that is to say, of very active drugs are prepared by percolation with Alcohol (70 p c), and in such a manner that the weight of the resulting tincture is equal to ten times the weight of the substance employed, in accordance with the *Brussels Convention*, 1902

Prescribing Notes—Most of the Tinctures mix readily with Water, but resinous Tinctures under similar circumstances require the addition of Mucilage of Gum Acacia, which is the best all-round emulsifying agent for this purpose It gives good results with all the Tinctures except Compound Tincture of Benzoin, which is very difficult to diffuse in Water, neither Mucilage of Gum Acacia nor Mucilage of Tragacanth, by itself, gives a satisfactory emulsion with this Tincture, the best effect is obtained by the use of Compound Tragacanth Powder, 60 grains of which will diffuse 3 fl drms of Compound Tincture of Benzoin, in 3 fl oz of Water

The quantity of Mucilage required for resinous Tinctures will depend upon the proportion of Tincture to the Water or other aqueous fluid, 1 fl drms of Mucilage of Gum Acacia is sufficient for 1 fl drms of the following Tinctures in 1 fl oz of Water —Benzoin, Cubebæ, Ammoniated Guaracum, or Tolu The following Tinctures require only about half this quantity —Asafetida, Cannabis Indica, Jalap, Myrrh, or Sumbul When Tincture of Hydrastis or Tincture of Podophyllum is prescribed with an aqueous solution of mineral salts, it is better to

add Mucilage of Gum Acacia The Mucilage should always be diluted with 3 or 4 times its bulk of Water before adding the Tincture

Mucilage of Tragacanth is also useful for the purpose of diffusing the Resin of the Tinctures, especially for Tincture of Jalap, and Tincture of Cannabis Indica with salts

Quinine is sometimes prescribed in mixtures under conditions which cause a precipitation of the alkaloid itself, or one of its sparingly soluble salts, in such cases the addition of 2 or 3 fl drms of Mucilage of Gum Acacia to the 6 or 8 oz mixtures will prevent the aggregation of the precipitate which would otherwise occur.

Not Official
TINOSPORA

The dried Stem of *Tinospora cordifolia*, Miers, is official in the *Ind* and *Col* *Add* for India and the Eastern Colonies, also *Infusum Tinosporæ* (1 in 10), dose $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 cc, *Liquor Tinosporæ Concentratus* (1 in 2), dose 30 to 60 minims = 1 8 to 3 6 cc, and *Tinctura Tinosporæ* (1 in 5), dose 30 to 60 minims = 1 8 to 3 6 cc.

Not Official.
TODDALIA

The dried Root-bark of *Toddalia aculeata*, Pers., is official in the *Ind* and *Col Add* for India and the Eastern Colonies, also *Infusum Toddaliæ* (1 in 10), dose 1 to 2 fl oz = 14 to 28 4 c c, *Liquor Toddaliæ Concentratus* (1 in 2), dose 30 to 60 minims = 1.8 to 3.6 c c

TRAGACANTHA.

TRAGACANTH

FR, GOMME ADRAGANTE, GER, TRAGANTH ITAL, GOYMA ADRAGANTE, SPAN,
GOMA TRAGACANTO

Thin, translucent, white, or pale yellowish-white odourless flaky shreds or filaments, possessing a somewhat horny appearance. It is a gummy exudation obtained by incision from *Astragalus gummifer*, Labill., and some other species of *Astragalus*.

The characteristic of the Syrian Tiagacanth is the form of ribbon-like flakes in which it occurs, and its comparative freedom from Starch

Pure *Triagacanth* gives a blue coloration with Iodine, varying in depth in different samples, but in any case it is much too faint to be confounded with added Starch.

Medicinal Properties—Demulcent Used for the suspension of heavy insoluble powders in liquids, 10 grains of the Compound Powder of Tragacanth are used for each fl oz of Water.

1 part of Tragacanth gives more viscosity to Water than 25 parts of Gum Acacia

The **Mucilage** is contained in **Lotio Hydragryi Nigra**

Not Official.—Bassorin, Gelanthum, and Glucantha.

Foreign Pharmacopœias—Official in Belg, Dan, Dutch, Fr, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Descriptive Notes—Tragacanth is found in various forms in commerce. The most valuable consists of semi-translucent thin flakes, known in commerce as Syrian Tragacanth, but imported from Persia. This is the official kind. It is 1 to 3 in (25 to 75 mm) or more in length and $\frac{1}{4}$ to 1 in (6 to 12 mm) in width, more or less contorted, white, translucent, horny, not easily broken but slightly flexible. The *P G* gives the dimensions as at least 0.5 cm broad and 1 to 3 mm thick, and its appearance as white and translucent. It is from 1 to 3 mm thick and is more easily pulverisable by a heat of 50° C (122° F), *USP*. Unlike Gum Arabic, it contains Starch grains. The kind known as Smyrna Tragacanth, which is next in quality, is more opaque and occurs in shorter, rather thicker flakes, which, owing to their greater opacity, have a faint, yellowish-white appearance. Small, slender strips are known as Vermicelli Tragacanth. Large, thicker pieces with a reddish tinge are known in trade as Gum Dragon and are used by shoemakers for smoothing rough leather, and for other technical purposes. A variety in small rounded pieces is known as Hog Gum or Caramania Gum, and is used for adulterating small Smyrna Tragacanth. It appears to be derived from *Astragalus Herutensis*, Bunge.

Tests—Tragacanth is sparingly soluble in Water, but swells up into a gelatinous mass which gives a violet or blue coloration with Iodine Solution, varying in depth in different samples, but in any case the coloration is much too faint to be confounded with that of added Starch. The *BP* states that it may be tinted violet or blue by Iodine Solution. The *USP* and the *P G* state that Tragacanth treated with 50 parts of Water swells up and gradually forms a cloudy gelatinous mass, which, when warmed on a water-bath with Solution of Sodium Hydroxide assumes a yellow coloration. In performing the test the *P G* employs powdered Tragacanth, the *USP* Tragacanth. The *USP* states that this gelatinous mass is tinged blue on the addition of Iodine T S, the *P G* that when the Tragacanth mucilage is diluted with Water, and the fluid filtered, Iodine Solution added to the residue on the filter produces a blackish-blue coloration, the filtered fluid is not coloured blue by Iodine Solution. The *USP* states that the addition of Alcohol (94.9 p c) to the fluid portion causes a precipitate, but the liquid is not coloured blue by Iodine T S. Tragacanth leaves when ignited with free access of air from 2 to 3 p c of ash, and 4 p c is rarely exceeded.

Preparations

GLYCERINUM TRAGACANTHÆ GLYCERIN OF TRAGACANTH

Tragacanth, in powder, $\frac{1}{2}$, Glycerin, $1\frac{1}{2}$, Distilled Water, $\frac{1}{2}$

Used as a pill excipient, but the following is better for that purpose—Tragacanth, in powder, 1, Glycerin, 6, rub together and keep for 2 or 3 days before use to allow it to stiffen.

'Diluted Glucose' is better than either

Official in Dutch, Tragacanth 1, Glycerin 9

MUCILAGO TRAGACANTHÆ. MUCILAGE OF TRAGACANTH

Mix 60 grains of *Tragacanth* in powder, with 2 fl oz of Alcohol (90 p c), in a bottle, add Distilled Water *q s* to form 10 fl oz and shake immediately (1 in 74)

Foreign Pharmacopœias—Official in Dutch, 1 in 50, Fr, 1 in 10, Ital and Port, 1 in 10, also 1 in 100, Mex, 1 in 20, Jap, Tragacanth 1, Glycerin 5, Tepid Distilled Water 94, Russ, Tragacanth 4, Acacia 1, Water 500, U S, Tragacanth 6, Glycerin 18, Water *q s* to make 100

PULVIS TRAGACANTHÆ COMPOSITUS COMPOUND POWDER OF TRAGACANTH

Tragacanth, 1, Gum Acacia, 1, Starch, 1, Refined Sugar, 3 (1 in 6)

Dose.—20 to 60 grains = 1 3 to 4 grammes

Not Official

BASSORIN—Gum Tragacanth 5, Glycerin 2, Water 93—*St John's*

It is also known as **Linimentum Exsiccans**

It can be medicated with 5 p c of Salicylic Acid, Hydronaphthol, or Thionin, with 10 p c of Acid Boric, or with 30 p c of Ichthyol, Resorcin or Precipitated Sulphur

Under the name of **Bassorin**, which is properly applied to the insoluble part of Tragacanth, there was introduced from the Continent a few years ago an ointment-basis made by mixing 1 part of powdered Tragacanth with spirit to wet it, then adding 50 parts of Glycerin (by weight) and heating until clear Martindale quotes the following formula—Tragacanth 5, Glycerin 2, Spirit 10, Water to 100 In the spirit contained in a wide jar the Tragacanth and add the Water, then add quickly the (as much Water, and shake well—*Pharm Form*

This has been incorporated in the *B P C*

GELANTHUM (*Unna*)—A firm basis used in dermatology, consisting of Gelatin, Tragacanth, Glycerin and Water

GLUCANTHA—Tragacanth, in powder, 240 grains, Water, 240 minims, Syrup of Glucose, 2 oz Pill Excipient—*Guy's*,

Not Official.**TRIFOLIUM****CLOVER**

A fluid extract is made from the dried *Trifolium pratense*, L, and from this a syrup, a teaspoonful of which 3 or 4 times a day is serviceable in *Menyanthes trifoliata*, L

Not Official**TRIMETHYLAMINA****TRIMETHYLAMINE**

C_2H_7N , eq 58.67

As supplied in commerce it is a colourless or pale yellow transparent solution, possessing a strong distinctive odour and a strongly alkaline reaction. It occurs somewhat frequently in both the animal and vegetable kingdoms. It is a constituent of the herring-bone, and has been detected in urine, unputrid blood of the calf and other animal fluids. It has been detected in America root, the blossoms of the Pear, Whitethorn, Hawthorn and

Wild Cherry The Trimethylamine of Ergot is probably a decomposition product of Choline. It is also a product of the dry distillation of certain alkaloids, Wood, etc., and especially of the masses or residues left after the distillation of the spirit from fermented beetroot molasses.

Propylamine is sometimes used as a synonym for Trimethylamine, but although isomeric with this substance its use as a synonym is not justified.

It is miscible with Water and with Alcohol (90 p.c.). It forms crystallisable salts. The Hydrochloride is the one chiefly used in medicine. Pure Trimethylamine is a gas at ordinary temperatures.

Tests—Trimethylamine has a sp. gr. at 0° C (32° F) of 0.673. It boils between 9° and 10° C (48.2° and 50° F). It is inflammable. It mixes readily with Water, forming a solution which is strongly alkaline in reaction towards Litmus paper. It combines with Carbon Bisulphide with evolution of heat. A glass rod moistened with Trimethylamine evolves white fumes when brought into contact with the vapour of Hydrochloric Acid. It combines with acids to form salts which are mostly crystallisable. Trimethylamine may be distinguished from primary and secondary Methylamines by its negative reaction with Alcohol Potash and Chloroform, that is to say, it does not evolve the characteristic and highly disagreeable odour of the trimethylamine or isomethylamine when boiled with Alcoholic Potassium Hydroxide, and Chloroform, by yielding no reaction when mixed with $\frac{1}{2}$ times its weight of Ethyl Oxalate (previously dried over Calcium Chloride), and by not effording volatile Nitrosamine when distilled with Nitrous Acid, and by its solution in excess of Hydrochloric Acid being precipitated by Potassium Ferriocyanide. When neutralised with Acetic Acid, the aqueous solution of Trimethylamine yields with Mercuric Chloride Solution a white precipitate. It gives with Iodine and with Iodo Potassium Iodide (Wagner's) Solution a yellow precipitate with Tannic Acid Solution a white precipitate, with Potassium Mercuric Iodide (Mayer's) Solution a white precipitate, and with Phosphomolybdic Acid a pale yellow precipitate. It may be determined by titration with Normal Volumetric Sulphuric or Hydrochloric Acid Solution, using Litmus Solution as an indicator of neutrality. 1 c.c. of the Normal Volumetric Acid Solution corresponds to 0.05867 gramme of absolute Trimethylamine.

TRIMETHYLAMINÆ HYDROCHLORIDUM—Translucent, colourless, very deliquescent crystals, possessing a strong distinctive odour, soluble in Water and in Alcohol (90 p.c.). It should be kept in well stoppered bottles of a dark amber tint in a cool atmosphere and protected as far as possible from contact with the air, as it is very deliquescent. It has been used in rheumatism and gout.

Dose—1 to 5 grains = 0.06 to 0.32 gramme.

Tests—Trimethylamine Hydrochloride dissolves readily in Water, forming a solution which has a neutral reaction towards Litmus paper. When mixed with Sodium Hydroxide Solution it evolves a powerful distinctive odour of Trimethylamine, the base separated from the salt should answer the tests distinctive of Trimethylamine given under that heading. It should dissolve in 10 parts of Absolute Alcohol, indicating the absence of Ammonium Chloride. The odour evolved on mixing it with Sodium Hydroxide Solution should possess the distinctive odour of Trimethylamine, and not an ammoniacal one. It should leave no weighable residue when ignited with free access of air.

Not Official TRITICUM

COUCH GRASS

The Rhizome of *Agropyrum repens*, Beauv., gathered in the spring and deprived of the rootlets.

Under the title **Agropyrum**, it is official, together with a **Liquid Extract** (1 in 1), in the *Ind* and *Col Add* for Australia, the Eastern and North American Colonies.

Medicinal Properties—Diuretic, and urinary sodative in cystitis and gonorrhœa

Official in U S, Austr, Belg and Swiss (*Rhizoma Graminis*), Fr (*Chien-dent*), Mex and Port (*Gramma Francoza*)

DECOCTUM TRITICI—Triticum, cut small, 1 oz, Water, 20 fl oz, boil 10 minutes, and strain when cold

Dose—4 to 8 fl oz = 113.6 to 227.2 cc 3 times a day

Fr, Tisane 1 in 50

A corresponding **Decoctum Agropyri**, dose $\frac{1}{2}$ to 2 fl oz = 14.2 to 54.8 cc, is *Ind* and *Col Add* for Australia, the Eastern and North American Colonies

EXTRACTUM TRITICI LIQUIDUM—Triticum, in No 20 powder, 10, Water until exhausted, evaporate the percolate to 15, and add 5 of set aside for 48 hours, filter, and make up to 20 with a mixture Rectified Spirit 1

Dose—1 to 6 fl drms = 3.6 to 21.3 cc

More easily prepared, and without heat (which is very detrimental to the Extract), by percolation with the above diluted Alcohol, so as to obtain 20 of finished product from 10 of the drug

Fluidextractum Tritici—Percolate 100 of Triticum with boiling Water until exhausted, evaporate the percolate to 75, and having added to it 25 of Alcohol (95 p c), mix well and set it aside for 48 hours, then filter the liquid and add sufficient of a mixture of Alcohol (95 p c) 1 and Water 3, to make 100. Average dose—2 fl drms = 7.1 cc—*U S P*

An extract is Official in Austr, Belg, Fr and Mex

TROCHISCI.

There are several lozenges in the Pharmacopœia They are made with four different bases

The **Simple Basis** consists of 496 of finely powdered Refined Sugar and 194 of Powdered Gum Acacia, made into a paste of 35½ of Mucilage of Gum Acacia and a small quantity of Distilled Water

Rose Basis is similar to the above, omitting 17½ of the Mucilage of Gum Acacia, and employing official Rose Water for making the paste

The **Tolu Basis** is similar to the Simple Basis, substituting 10½ of Tincture of Balsam of Tolu and 10½ of Distilled Water for a portion of the Sugar

Fruit Basis is similar to the Simple Basis, substituting 56½ of Black Currant Paste for the same quantity of Sugar

Compressed Lozenges—The general method is to granulate the mixture of medicament and Gum, by means of Theobroma Emulsion (p 1191) and highly compress the dried granules (*C D '03*, n 231) The advantage of avoiding the application of heat is obvious in the case of volatile substances, such as Phenol and essential Oils—*F J '03*, n 158

Not Official

TYLOPHORÆ FOLIA.

The dried Leaves of *Tylophora asiamatica* dose $\frac{1}{2}$ to 2 grains = 0.016 to 0.13 gramme as an expectorant, 15 to 30 grains = 1 to 2 grammes as an emetic, are official in the *Ind* and *Col Add* for India and the Eastern Colonies,

Not Official
U L E X I N E

Syn—CYTISINE

A crystalline alkaloid prepared from *Ulex Europæus*, L., the common gorse or furze

Solubility—Freely soluble in Water and Chloroform, insoluble in pure Ether

The Nitrate, Hydrochloride, and Hydrobromide are crystalline salts readily soluble in Water

Medicinal Properties—Diuretic useful in cardiac dropsy

Dose— $\frac{1}{20}$ to $\frac{1}{15}$ grm = 0.0032 to 0.0012 grmme dissolved in 60 minims of Water

Ulexine temporarily masks the action of Strychnine —*T G* '87, 280, 690

Not Official
U L M U S

Under this title the dried inner bark of *Ulmus campestris*, L., was official in *B P* '64 and '67, the dried bark of *Ulmus fulva*, Mich., deprived of its periderm, is official in the *U S P*. The value of both the barks depends upon the mucilage which they contain, that of the *Ulmus fulva* is stated also to preserve fatty substances from becoming rancid

Decoctum Ulmi (*B P* '67), Elm Bark 1, Water 8, boil for 10 minutes, strain and make up to 8, dose 2 to 4 fl oz = 56.8 to 113.6 cc 3 or 4 times daily

This has been incorporated in the *B P C*

Mucilago Ulmi (*U S*), 6 of Slippery Elm (*Ulmus fulva*) in 100 of Water, digest in a covered vessel, on a water bath for one hour, and strain

This has been incorporated in the *B P C*

U N G U E N T A.

For the preparation of the Ointments of the British Pharmacopœia, various bases are used, *eg*, Soft Paraffin, Hard Paraffin, a mixture of Hard and Soft Paraffins, Lard, Benzoyated Lard, Beeswax and Lanolin

In the case of the ointments containing alkaloids, Oleic Acid is used with the object of dissolving the alkaloid. In India and the Colonies, when the ointment would be too soft, owing to the warmer climate, indurated Lard, prepared Suet or Beeswax may be employed for the purpose of stiffening the ointment, provided such admixture does not affect the proportion of active ingredient

Eye Ointments—The basis for these is neutral yellow Soft Paraffin, which has been melted and strained through fine muslin. The medicament in very fine powder should be first rubbed with a small portion of the Paraffin, and in the case of alkaloids the Paraffin may be warmed (not above 50° C) until solution is effected —*St Thomas's*

This has been incorporated in the *B P C*

Ointments appear in the Foreign Pharmacopœias, under the following generic titles —

Austr, Belg, Dan, Dutch, Ger, Hung, Jap, Norw, Russ, Swed, Swiss and *U S*, Unguenta, Fr (*Pommades*), Ital (*Pomato*), Mex, Port and Span (*Unguento*)

Not Official.

URANIUM NITRATE

Pale yellow, rhombic crystals, readily soluble in Water. It should be kept in well-stoppered bottles and protected as far as possible from the light. The *B P Appendix* describes it as the crystals of pure Uranium Nitrate of commerce. Used in diabetes.—*B M J* '95, ii 467, '97, ii 1014, *P* 1257

Dose—1 to 5 grains = 0.06 to 0.32 gramme

Tests—Uranium Nitrate when heated melts, loses its Water of crystallisation, and when more strongly heated loses also Nitric Acid. It dissolves readily in Water, forming a clear solution which is acid in reaction towards blue Litmus paper. This aqueous solution affords with Ammonium Hydrosulphide Solution a chocolate brown precipitate insoluble in excess of the reagent. Ammonium, Potassium or Sodium Hydroxide Solution produces a yellow precipitate insoluble in excess of the reagent. In the presence of Tartaric Acid these reagents do not produce a precipitate, the precipitate produced by Ammonium Hydroxide Solution is soluble in a solution of Ammonium Carbonate. Ammonium, Potassium or Sodium Carbonate Solution yields a light yellow precipitate soluble in excess of the reagents. Potassium Ferrocyanide Solution produces a reddish-brown precipitate in sufficiently concentrated solution, or a reddish-brown coloration even in highly diluted solutions. Potassium Ferrocyanide Solution produces no change, Sodium Acetate Solution, more particularly in the presence of Sodium Acetate and Sodium Chloride, produces a whitish precipitate. A standard solution of Uranium Nitrate is used for the determination of Phosphoric Acid, Potassium Ferrocyanide Solution being employed as an indicator.

URANIUM SALICYLATE—A pale yellowish green crystalline salt, seems (*L* '05, i 387) to be better tolerated in cancer than either the Acetate or Nitrate. **Dose**, 5 to 20 grains = 0.32 to 1.3 grammes

The Uranium compounds have lately received a very considerable amount of attention, the metal first gave rise to a suspicion of the existence of a radioactive property in elements, and this suspicion was followed by M et Madame Curie's discovery of the radio-active element, Radium, in pitchblende.

RADIUM—A lengthy and intricate process for the separation of this radio-active element has been fully recorded by its discoverers, M et Madame Curie, in the *Annales de Chimie et de Physique*, to the Faculté des Sciences de Paris, and reprinted in series in the *Chemical News*, and summarised, *L* '03, ii 966. The salt chiefly used in medicine is the Radium Bromide, which is usually supplied in capsules containing 0.005 mg (about $\frac{1}{40}$ grain). It is a white salt, but gradually becomes coloured.

The peculiar action of the rays on tissues has been utilised in the treatment of carcinomatous and sarcomatous growths, in epithelioma, psoriasis and lupus.—*L* '03, ii 271, 927, 966, 1388, *B M J E* '03, ii 31

Applied to the skin for 20 to 40 minutes or longer in lupus, rodent ulcer and superficial epitheliomata.—*B M J* '03, ii 199

Treatment of consumption by the rays from Radium and Thorium.—*B M J* '03, ii 197

It has been discovered in the waters of Bath and Burton. The deposits from these mineral waters were estimated each to contain about the same amount, the amount in the deposit being relatively much greater than that in solution.—*B M J* '04, i 797

17 cases of cancer treated by the emanations of 30 mg enclosed in a vulcanite capsule covered with talc. It appeared that the emanations from Radium can only act upon the rapidly growing cells, and that the older cells, which are surrounded by fibrous tissue, are less and less easily affected and are therefore in excess of fibrous tissue the cells are not at all affected.—*L* '04, i 1047.

Recent cases obtained in asthma by a 20 minutes application on the chest, followed by a 25 and 30 minutes' application respectively on two successive days.—*B M J* '04, ii 1234

In dermatology 10 mg of Radium Bromide applied in several sittings daily.—*B M J E* '05, ii 15

In the treatment of rabies (*B M J* '05, ii 36), animals were inoculated with the virus and exposed for some days to the action of Radium. Controls inoculated with virus of equal strength and not submitted to the same treatment all died.

A record of 9 cases of cancer of the oesophagus treated with Radium. In 6 cases the treatment was so far successful as to cause some widening of the stricture. In the other 3 no improvement took place. 3 to 1 hour's application is made daily or every other day for several weeks.—*B M J* '05, ii 92.

6 cases of malignant tumour, 5 of which were carcinomata, and 1 of melano sarcoma, treated by 10 mg of the Bromide. In no case did the treatment prove of any value. Not recommended for cancer of any kind. In operable cases the knife yields infinitely more promise, and in inoperable cases Radium only does harm. All the lupus cases were cured.—*B M J E* '05, i 39.

1 mg of the Bromide enclosed in a thin glass tube of 3 cm length and 2 mm diameter, in the treatment of granulation of the conjunctiva.—*B M J E* '05, i 43. The exposure was carried out daily for 10 to 15 minutes, and resulted in cure.

In the treatment of rodent ulcer. A tube containing 5 mg of the Bromide applied by tying the Radium tube between the ulcer and a layer of gutta serena tissue, the durations of the applications averaging 20 minutes. Whether the results will be as permanent as after the usual treatment has yet to be proved (*B M J* '05, ii 9), but no one seeing the new skin can have any doubt of the greater perfection of cosmetic effect over any treatment hitherto known.

5 mg Radium Sulphate of 500,000 units attached with enamel varnish to a plate of Copper 1 in square, applied for 30 minutes to each lobe of a trilobate tumour affecting the upper eyelid, the exposure being repeated 3 days afterwards. The tumour had melted away, leaving only a small ulcer.—*L* '05, ii 548.

A method of coating instruments, celluloid rods, discs, etc, with Radium (*L* '05, ii 545), a salt of the latter being dissolved in a suitable volatile solvent tinted with an aniline dye and the instrument dipped in.

Thorium Nitrate, Thorium Lactate, and Thorium Salicylate are salts of the rare metal, Thorium, which have been introduced and which have found more or less use commercially.

Not Official

UREA

CARBAMINE, CARBONYLAMIDE

$\text{CH}_4\text{N}_2\text{O}$, eq 59.67

Colourless, transparent, almost odourless, somewhat hygroscopic, prismatic crystals, possessing a cool, saline taste.

Solubility—1 in 1 of Water, 1 in 7 Alcohol (90 p.c.)

Introduced as a diuretic, it can dissolve uric acid calculi.—*L* '01, i 694, 1672, '01, ii 1567, 1709, '02, i 548, '02, ii 1383, 1456, '03, ii 1017, *B M J* '02, ii 1235.

20 grains 3 times a day gradually increased to 120 grains 3 times daily, combined with the application of the X rays, in lupus vulgaris.—*L* '02, i 659.

It is stated to possess the power of dissolving coagulated proteids.—*L* '02, ii 527, *P J* '03, i 385.

Dose—20 to 60 grains = 1.3 to 4 grammes, 3 or 4 times daily.

Hypodermically it may be given in 40 grain doses dissolved in 4 fl. drms sterilised Water.

Tests—Urea melts at about 132.5° C (270.5° F), and at a temperature of 150° to 160° C (302° to 320° F) it is decomposed with the evolution of Ammonia and formation of Biuret. It dissolves readily in Water, forming a solution which is neutral in reaction towards Litmus paper. At the ordinary temperature the solution has no tendency to change, but on boiling it is decomposed with the formation of Ammonium Cyanide. Urea when heated in a test tube melts, and then evolves Ammonia, when fused with Potassium or Sodium Hydroxide or

ignited with Soda-Lime, Ammonia is also evolved, recognised by its distinctive odour and by its reaction on a piece of moistened red Litmus paper which it turns blue. When heated for some time to a temperature not exceeding 160°C (320°F), cooled, the residue dissolved in Water, mixed with Sodium Hydroxide Solution and then with diluted Cupric Sulphate Solution, a violet or red coloration is produced, this reaction is known as the Biuret test. When moistened with concentrated solution of Furfural, and a drop of Hydrochloric Acid (sp gr 1.1) a fine violet coloration is produced. An aqueous solution when heated with Silver Nitrate affords a white precipitate of Silver Cyanide. Urea is not precipitated by Mercuric Chloride Solution, nor by a solution of Mercuric Acetate. It is not precipitated by Tannic Acid Solution, by Potassium-mercuric Iodide (Mayer's) Solution, by Iodo-potassium Iodide Solution, Phosphoric Acid Solution, nor the other general reagents for Urea. It yields no reaction with either neutral or basic Lead Acetate Solution, it does not reduce Fehling's Solution even on boiling. When mixed with Sodium Hypobromite Solution it evolves Nitrogen, and this reaction is utilised for its determination when necessary, the absence of substances similarly evolving Nitrogen on treatment with Hypobromite being first assured. When ignited with free access of air it should leave no weighable residue.

UROL (Urea Quinate) —Large, colourless, prismatic crystals, having an acid, bitter taste, readily soluble in Water and in Alcohol (90 p c).

It has been recommended in the Uric Acid diathesis.

VERONAL Diethyl-malonyl Urea $\text{C}_2\text{H}_5\text{C}_2\text{H}_4\text{C}_4\text{N}_2\text{H}_2\text{O}_8$, eq 182.80 —Colourless, odourless crystals, or a white, crystalline powder, possessing a faintly bitter taste.

Solubility —1 in 160 of Water, 1 in $8\frac{1}{2}$ of Alcohol (90 p c).

A hypnotic. It is given a high place (*B M J E* '04, ii 96) as a sleep-producing agent, the effect being chiefly sedative and of little value where there is pain. Although a good hypnotic (*B M J* '04, ii 1679), it seems to take time to act, and to have a cumulative action, unfavourable results following the administration of 8 doses of 10 grains given at intervals of 1 hour. In a case of mental excitement (*B M J* '04, ii 1784), where 10 grains thrice daily had been taken for a week toxic symptoms followed. 2 doses of 10 grains each caused urticaria, lasting 8 days, and in the other case local cedema lasting a week —*B M J* '04, ii 1736.

A most satisfactory hypnotic, very seldom, except in mental cases, will more than 7 or 8 grains be required for a dose, writers differ very widely as to the dose, best given in hot fluid —*F T* '07, 73.

It acts with comparative certainty in small doses and without deleterious effects. The best of the non-Chlorine hypnotics, and ranks with Chloral —*B M J* '05, ii 250.

Best given periodically, and often varied. The smallest effective dose should be used in the commencement, and the drug removed from the system at the first opportunity —*M P* '05, ii 563.

Combining anodynes with hypnotics for administration at night is pain pointed out (*B M J* '05, ii 1008). A small dose of Aspirin added to Tronal or Veronal will produce sleep under many circumstances where the hypnotic alone will fail.

Acts mildly and produces a sleep which is very like that of nature. It fails when there is much pain. Of the unpleasant side effects are mentioned the production of rashes and the diuretic action —*B M J E* '05, ii 4.

Appears to combine certainty of action with the advantages of inducing sleep in such small doses (5 to 10 grains) as have hitherto proved efficacious only in the Chlorine compounds —*B M J* '05, ii 1005.

Fatal case of Veronal poisoning —*L* '05, ii 234. $\frac{1}{2}$ oz taken between a Thursday and a Saturday morning. Attention called to the unrestricted sale of such a quantity as 1 oz of the drug to a private person.

Ought to be administered always with great caution in small doses, and especially care ought to be taken in cases of renal insufficiency —*J J E* '05, ii 63, *B M J* '07, i, 269.

Dose —5 to 20 grains = 0.32 to 1.3 grammes

Prescribing Notes—*It can be made into pills containing 5 grains each, with $\frac{1}{2}$ of its weight of 'Diluted Glucose' It can also be dispensed in cachets*

Tablets are supplied containing $7\frac{1}{2}$ grains = 0.5 grammes in each

Official in Swiss, Acidum Diethylbarbituricum

Tests—Veronal melts at 191°C (375°F). It sublimes without residue, except possibly a faint trace of Carbon. It dissolves sparingly in Water, forming a neutral solution, but is more readily soluble in Alcohol (90 p.c.). The saturated aqueous solution acidified with Nitric Acid yields on the addition of Millon's reagent (1 part by weight of metallic Mercury dissolved in the cold in 1 part by weight of fuming Nitric Acid, the solution diluted with 2 parts of Distilled Water, and filtered) a white colourless precipitate. 0.2 of a gramme of Veronal when fused with Potassium Hydroxide evolves Ammonia, recognisable by its distinctive odour and by its action upon moistened red Litmus paper, if the cooled residue be acidified with Diluted Sulphuric Acid, Carbon Dioxide is evolved and a characteristic fatty odour is developed. 1 gramme when ignited with free access of air should leave no weighable residue.

ELIXIR DIETHYLBARBITURIC ACID—Diethylbarbituric Acid (Veronal), 18 grammes, Compound Tincture of Vanilla (NF), 16 c.c., Alcohol, 175 c.c., Glycerin, *qs* to make 500 c.c. Dissolve the Diethylbarbituric Acid in the Alcohol, add the Compound Tincture of Vanilla, and enough Glycerin to make 500 c.c.—*C D '03, 11 521*

Bromural (Urea Monobromine Isovalerianate)—In white platelets having a slightly bitter taste, soluble in hot Water, Ether, Alcohol and the alkalis. The dose is 5 to 10 grains = 0.32 to 0.65 gramme, introduced as a hypnotic.—*B M J E '07, 1 75*

Not Official URETHANE

ETHYL CARBAMATE ETHYL URETHANE CARBAMIC ACID ETHYL ESTER

$\text{C}_3\text{H}_7\text{NO}_2$, eq 88.43

Colourless, prismatic, odourless crystals or scales, with a peculiar cool taste. Urethane is official in the *U S P* under the title of *Æthylis Carbamas*. It may be prepared by the action of Ethyl Alcohol upon Urea or one of its salts. It should be kept in well stoppered glass bottles, preferably of a dark amber tint.

Solubility—1 in 2 of Water, 1 in 1 of Alcohol (90 p.c.), 2 in 3 of Ether

Medicinal Properties—Hypnotic, without anodyne properties.

Possesses a slightly irritant action.—*L '99, 11 72*

Was good as a hypnotic, but it had to be used in very large quantities.—*B M J '05, 11 250*

Is uncertain and weak in action.—*B M J '05, 11 1005*

Dose—15 to 30 grains = 1 to 2 grammes

Official in Spain, Swiss and Mex. (Uretano)

Tests—Urethane melts at about 48°C (118°F). The *U S P* states 47 to 50°C (117 to 122°F), it boils at about 172°C (341°F). At a higher temperature it is decomposed. When mixed with 5 times its weight of Sulphuric Acid and gently heated it is decomposed with the evolution of Carbon Dioxide. When warmed with Sodium Hydroxide Solution (15 p.c.) the distinctive odour of Ammonia is evolved, and a piece of red Litmus paper suspended in the mouth of the tube is rendered blue. 0.5 of a gramme dissolved in 5 c.c. of Water, containing in solution 1 gramme of dry Sodium Carbonate yields when the solution is warmed with the addition of Iodine a yellow crystalline precipitate of Iodoform when the solution cools. The 10 p.c. aqueous solution should not afford a turbidity on the addition of Silver Nitrate Solution, indicating the absence of Chlorides. 2 c.c. of a 10 p.c. aqueous solution mixed with 2 c.c. of cold concentrated Sulphuric

Acid, the liquids being mixed, should not yield a brownish tint at the junction of the two liquids. Separate solutions of 1 c.c. of Ferrous Sulphate Solution, indicating the absence of Nitrates. Separate solutions of 1 gramme of Urethane dissolved in 1 c.c. of Water should neither afford a crystalline precipitate on the addition of 1 c.c. of Nitric Acid, nor on the addition of Mercuric Nitrate Solution, nor on the addition of Oxalic Acid Solution, indicating the absence of Urea or Carbamide. 1 gramme when heated with free access of air should leave no weighable residue, indicating the absence of mineral impurities.

HEDONAL (Methyl propyl-carbinol-methane).—Colourless crystals, or as a white crystalline powder, slightly soluble in cold Water, but more readily in hot Water.

Introduced as a hypnotic. Stated (*B M J E* '05, 134) to have been given as a hypnotic to supplement Chloroform anaesthesia in doses of 30 grains from 1½ to 1 hour before operation, small quantities of Chloroform then sufficing to produce anaesthesia.

Dose—15 to 30 grains = 1 to 2 grammes, in cachet.

Somnal.—Stated to contain Urethane and Chloral Hydrate, was introduced as a hypnotic, in doses of 30 grains.

Phenyl-Urethane (Euphorin).—A white crystalline powder, only sparingly soluble in Water, soluble in Alcohol (90 p.c.) and in Ether. It should be free from the light. A powerful analgesic, but like some other powerful tends to interfere with the respiratory processes and to weaken the heart. It has proved of special service in the pain of orchitis. Dose—1 to 5 grains = 0.06 to 0.32 gramme.—*B M J* '98, 11 1055.

UVÆ URSI FOLIA.

BEARBERRY LEAVES

FR, BUSSESOLE, GER, BARENTRAUBENLATTEP, ITAL, UVA URSINA,
SPAN, GAYUBA

The dried Leaves of *Uva-ursi*, Spengel

Contains a crystallisable glucoside, Arbutin, soluble in Water and Alcohol (90 p.c.), dose, 1 to 15 grains.

Medicinal Properties.—Astringent and diuretic, it is a disinfectant to the urinary mucous membrane, and is valuable in inflammation of the bladder and urethra.

Official Preparation.—Infusum Uvæ Ursi

Not Official.—Infusum Uvæ Ursi Concentratum

Official in Aust, Belg, Dan, Dutch, Fr (Bussesole), Ger, Ital, Jap, Mex (Gayaba del pais), Norw, Port (Uva Ursina), Russ, Swed, Swiss and U.S.

Descriptive Notes.—The leaves are about $\frac{3}{4}$ to 1 in long (19 to 25 mm) long and $\frac{1}{4}$ to $\frac{1}{2}$ in (6 to 9 mm) broad, obovate, rounded at the apex, and tapering below into a short leaf stalk, dark green and shining on the upper surface, with a network of depressed small veins, the under surface pale and reticulated with dark veins, the margin is entire and slightly revolute. The taste is astringent, and the odour faint and herbaceous. The leaves of *Vaccinium Vitis-Idæa*, L., bear some resemblance to Bearberry Leaves, and are stated to have been mixed with them, but can be easily distinguished by having dark dots on the under surface, by being coarsely toothed near the apex and more revolute at the margin.

The powdered leaves are characterised by the straight-walled

epidermal cells, the large stomata of the lower epidermis, short palisade cells, and the presence of tracheids and numerous serial prismatic crystals

Tests—Bearberry Leaves leave from 2 to 3 p c of ash.

Preparation

INFUSUM UVÆ URSI INFUSION OF BEARBERRY

Bearberry Leaves, bruised, 1, boiling Distilled Water, 20, infuse for 15 minutes and strain (1 in 20)

Dose— $\frac{1}{2}$ to 1 fl oz = 14 2 to 28 4 c c

In the 1864 Pharmacopœia the Leaves were not ordered to be bruised, *when bruised*, the infusion is stronger, but a large deposit forms in the strained fluid

Incompatibles—Iron salts, Lead salts, Silver Nitrate, vegetable alkaloids, Gelatin

Foreign Pharmacopœias—Official in Fr (Tisane), 1 in 100, Ital, 1 in 20 Decoction, U S has a fluid extract

Not Official

INFUSUM UVÆ URSI CONCENTRATUM—Bearberry Leaves, in No 20 powder, 40, Alcohol, (90 p c), 25, Dilute Chloroform Water (1 in 1000), *q s* to make 100 Prepare by the repercolation—*Far* and *Wright*, *P J* '06, 1 165 and '07, 1 621, *C D* '06, 1 252, and *I L P* 1907, 248

Dose— $\frac{1}{2}$ to 1 fl dm = 1 8 to 3 6 c c

This appears in the *B P C*

VALERIANÆ RHIZOMA.

VALERIAN RHIZOME

B P Syn—VALERIAN ROOT

FR, VALERIAN OFFICINALE, GER, BALDRIAN, ITAL, VALEPIANA, SPAN, VALERIANA

The dried elect Rhizome and Roots of *Valeriana officinalis*, L, collected in the autumn

That from wild plants growing on dry soil is preferred. It owes its properties to a volatile Oil and a volatile Acid, the salts of the latter (Valerianates) are not prepared from the root, but synthetically from Amylic Alcohol

The bulk of the Valerian root used in this country is of foreign growth, and should either be allowed or expressly prohibited in *B P*

Under the title **Valerianæ Indicæ Rhizoma**, the dried Rhizome and Rootlets of *Valeriana Wallichii*, DC, are official in the *Ind* and *Col* add for India and the Eastern Colonies

Medicinal Properties—It is a nerve stimulant and antispasmodic. Useful in hysteria, in functional nervous diseases associated with hysteria, and as an adjunct to tonics

The difference in physiological action between the juice and the dried root of Valerian is stated (*L* '05, 1 1396) to be due to oxidation of the active constituents during drying. The sedative and antispasmodic action of the fresh juice is very constant, and is not accompanied by any permanent stimulating action. Since the fresh juice owes its peculiar physiological properties to the undecomposed bornyl iso valerianate contained in the volatile Oil, it would appear to be more desirable to use the volatile Oil in preference to the other preparations of Valerian

Official Preparation—Tinctura Valerianæ Ammoniatæ

Not Official—Tinctura Valerianæ, Tinctura Valerianæ Ætherea, Oleum Valerianæ, Valyl, Acidum Valerianicum, Fluidextractum Valerianæ, Infusum Valerianæ, Infusum Valerianæ Concentratum

Foreign Pharmacopœias Official in all. An Extract, and a 1 in 20 Infusion are official in Ital. An Extract in Bel., Dutch, Fr., and Russ., a Fluid Extract in Dan., Mex. and U.S.

Descriptive Notes—Valerian Root varies much in quality and in price. A little is grown in this country at Chesterfield, where the form *sambucifolia*, Willd., appears to be the species cultivated, at Long Melford the more robust form *Mikanii*, Syme, is preferred, the latter yielding a rather larger and more odorous root. Valerian Root is also imported from Thuringia in Germany, Hungary, Belgium, and France, and rarely from Japan, under the name of Kesso, the Japanese plant is referred to the var. *latifolia*, Miq. The French root is generally rather pale, the Japanese is a dark brown with a scurfy surface, and is powerfully odorous, it is probably a distinct species.

Valerian Root consists of a short rootstock, $\frac{1}{2}$ to $\frac{1}{2}$ in (8 to 12 mm) in diameter and less than 1 in (25 mm) long, giving off numerous slender, brownish, brittle roots, 3 to 4 in (7.5 to 10 cm) long and about $\frac{1}{16}$ in (2.5 mm) in diameter, tapering into slender rootlets at the extremity, and whitish in transverse fracture. The rhizome is hard and hoary internally, but becomes hollow with transverse septa when old and occasionally exhibits a few lateral, short, horizontal branches. When fresh it is almost without smell, the valerianic odour being developed during the drying or by injury to the surface. Under the microscope the characteristic features are the hypoderm cells with undulated walls, the abundance of small rounded or muller-shaped starch grains, the oil drops in the cortical cells, and the porous sclerenchymatous cells of the rhizome.

Tests—Valerian Root yields from 8 to 10 p.c. of ash. The B.P. states that the odour developed in the process of drying is strong, characteristic, and disagreeable, the taste unpleasant, camphoraceous, and slightly bitter, the U.S.P. that the odour is peculiar, becoming stronger and more unpleasant on keeping the drug, the taste is camphoraceous and somewhat bitter.

Preparation

TINCTURA VALERIANÆ AMMONIATÆ AMMONIATED TINCTURE OF VALERIAN

Valerian Rhizome, in No. 40 powder, 4 oz., Oil of Nutmeg, 30 minims, Oil of Lemon, 20 minims, Solution of Ammonia, 2 fl. oz., Alcohol (60 p.c.), 18 fl. oz., by maceration.

Tests.—Tinct. Valerian Ammon. has a sp. gr. of 0.935 to 0.945, it contains about 3.5 p.c. w/v of total solids and about 53 p.c. w/v of Absolute Alcohol. When freshly prepared a measured quantity of 10 c.c. of the tincture requires about 4.3 c.c. of Normal Volumetric Sulphuric Acid Solution to neutralise the Ammonia, Methyl Orange,

or Cochineal Solution being employed as an indicator of neutrality
This corresponds to 0.72 p.c. w/v of absolute Ammonia

Dose — $\frac{1}{2}$ to 1 fl. drm. = 1.8 to 3.6 c.c.

Tinctura Valerianæ Ammoniata (U.S.)—Valerian, in No. 60 powder, 20, Aromatic Spirit of Ammonia, *q.s.* to make 100, by macero-percolation

Tinctura Valerianæ Indicæ Ammoniata—Indian Valerian, in No. 40 powder, 4 oz. Oil of Nutmeg, 30 minims, Oil of Lemon, 20 minims, Solution of Ammonia, 2 fl. oz. Alcohol (60 p.c.), 18 fl. oz., by maceration. Dose — $\frac{1}{2}$ to 1 fl. drm. = 1.8 to 3.6 c.c. It is official in the *Ind.* and *Col. Add.* for India and the Eastern Colonies

Not Official

FLUIDEXTRACTUM VALERIANÆ—100 of Valerian, in No. 40 powder, is first moistened with 30 of a mixture of Alcohol (95 p.c.) 75 and Water 25, macerated in a percolator for 48 hours, then exhaust, reserve the first 85 of percolate and evaporate the remainder at a temperature not exceeding 50° C. (122° F.) to a soft extract, dissolve this in the reserve portion, and make up with the menstruum to 100—*U.S.P.* Dose, 30 to 60 minims = 1.8 to 3.6 c.c.

This has been incorporated in the *B.P.C.*

This Fluid Extract evaporated to a firm extract constitutes **Extractum Valerianæ**—*B.P.C.*

INFUSUM VALERIANÆ—Valerian Rhizome, bruised, $\frac{1}{2}$, boiling Distilled Water, 10. Infuse in a covered vessel for 1 hour and strain—*B.P.* 1885

This is incorporated in the *B.P.C.*, infusing 15 minutes

INFUSUM VALERIANÆ CONCENTRATUM—Valerian Rhizome, in No. 20 powder, 40, Strong Solution of Ammonia, 0.3, Alcohol (90 p.c.), 25. Dilute Chloroform Water (1 in 1000), *q.s.* to make 100. Mix the powder with the Strong Solution of Ammonia and sufficient Chloroform Water to damp it evenly, set aside for 2 hours, and then submit to re-percolation—*Farr* and *Wright, P.J.* '06, 1.165 and '07, 1.622, *C.D.* '06, 1.252, and *E.B.P.* '07, 251

Dose — $\frac{1}{2}$ to 1 fl. drm. = 1.8 to 3.6 c.c.

This appears in the *B.P.C.*, using 20 of Valerian Rhizome instead of 40

TINCTURA VALERIANÆ—Percolate 1 of Valerian Rhizome, in No. 40 powder, with sufficient Alcohol (60 p.c.), to yield 8—*B.P.* 1885

Dose — 1 to 2 fl. drm. = 3.6 to 7.1 c.c.

This was included in the *B.P.C. Formulary* 1901

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Fr., Ger., Hung., Ital., Mex., Norw., Port., Russ., Swed., Swiss and U.S., 1 in 5, Jap., 1 in 10, Mex. and U.S. have also **Fluid Extract**. All by weight, except U.S.

Tests—Tincture of Valerian (*B.P.* '85) has a sp. gr. of 0.924 to 0.930, contains about 2.0 p.c. w/v of total solids and about 60.0 p.c. w/v of Absolute Alcohol

TINCTURA VALERIANÆ ÆTHEREA (Ger.)—Valerian, 1, Spirit of Ether, by weight, 5

Foreign Pharmacopœias—Official in Austr., Belg., Dan., Dutch, Ger., Hung., Jap., Norw., Span. and Swiss, 1 in 5, Mex., 1 and 5, Sp. Ether (sp. gr. 0.76), Russ., Valerian 1, Alcohol (90 p.c.) 4, Ether (0.725), 2. All by weight

Tests—Etheral Tincture of Valerian (*P.G.*) has a sp. gr. of about 0.815, and contains about 1.0 p.c. w/v of total solids

OLEUM VALERIANÆ—A yellow volatile Oil, sp. gr. 0.930 to 0.960. Dose — 2 to 5 minims = 0.12 to 0.3 c.c.

Foreign Pharmacopœias—Official in Austr., Belg., Hung. and Port.

VALYL (Diethylamide Valerianate)—An oily liquid, possessing a nauseous odour and taste. A sedative in nervous affections. Dose — 2 to 10 grains = 0.13 to 0.65 gramme

Best given in capsules—*B.M.J.E.* '02, 1.3

ACIDUM VALERIANICUM. Valerianic Acid, Valeric Acid $C_5H_{10}O$, eq 101.31—A transparent, colourless, or nearly colourless, oily liquid, possessing a strong distinctive disagreeable odour. It is used in the preparation of the Valerianate.

It should be kept in well stoppered glass bottles of a dark amber tint and in a cool place.

Official in Fr

Tests—Absolute Valeric Acid has a sp gr of 0.938 at $15^{\circ}C$ ($59^{\circ}F$). It boils about $175^{\circ}C$ ($347^{\circ}F$). Commercial Valeric Acid contains a varying proportion of the pure acid, it is recognised by its distinctive penetrating disagreeable odour. When warmed with a mixture of Sulphuric Acid and a little Ethyl or Amyl Alcohol it evolves a fragrant fruity odour. When neutralised with Ammonia and tested with Ferric Chloride Test a brownish-red precipitate is thrown down, when this precipitate is allowed to settle the supernatant liquid should be colourless, in the presence of Formic or Acetic Acid the supernatant liquid is coloured red. When concentrated Valeric Acid is agitated with Copper Acetate Solution, anhydrous Cupric Isovalerate separates in oily drops, which ultimately crystallise in greenish-blue monoclinic prisms, the reaction distinguishes Valeric Acid from Butyric Acid, the latter acid forming with a moderately concentrated Cupric Acetate Solution an immediate crystalline precipitate of Cupric Butyrate. The acid may be readily determined by direct titration with Normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. 1 cc of Normal Volumetric Sodium Hydroxide Solution corresponds to 0.10131 gramme of absolute Valeric Acid.

The acid should be completely volatile, and should leave no weighable residue.

Not Official

VANILLA.

The Fruit of *Vanilla planifolia*, Andr, chiefly used as a flavouring agent. The finest quality comes from Mexico, and large quantities also come from Bourbon. It owes its fragrance to **Vanillin**, which on oxidation yields **Vanillic Acid**. Some text books refer to them as the same substance, but this is not the case, Vanillic Acid is without odour and does not form a crystallisable compound with Sodium Bisulphite.

Foreign Pharmacopœias—Official in Austr, Belg, Fl, Ger, Jap, Mex, Swiss and U S. Swiss has Tincture 1 in 5, Fl and U S 1 in 10.

Descriptive Notes—Vanilla pods are the nearly ripe fruits of *Vanilla planifolia*, Andr, prepared by scalding, gradual fermentation and drying. After the curing process (*Lign. News*, vi, p 291, *P J* (4) vii, p 640) the pods are sorted out into various lengths so as to form bundles of uniform size. Mexican Vanilla is considered to be the most aromatic, the pods are 8 to 10 inches (20 to 25 cm) long, flattened, and about $\frac{3}{8}$ inch (9 mm) in diameter at the broadest part. The upper end tapers gradually to the point of attachment to the plant, and is curved and slightly twisted there. The longest pods obtain the highest price. When kept the pods become 'frosted' or covered with 'givre,' which consists of fine crystals of Vanillin. The value of Vanilla does not, however, depend upon the amount of Vanillin contained in the pods, but upon the aroma, which the artificial Vanillin cannot entirely replace.

VANILLIN ($C_8H_8O_2$, eq 140.92)—It is the Aldehyde of Methylprotocatechuic Acid and yields on oxidation Vanillic Acid ($C_8H_8O_3$). It is official in the *U S P*, and is stated to occur naturally in Vanilla or to be made artificially from several Ortho dihydrobenzoin derivatives. It is a white needle shaped the peculiar distinct odour and taste of Vanilla. It has an a

Foreign Pharmacopœias—Official in Austr, Belg, Fl, Ger, Jap, Mex, Swiss and U S.

Tests—Vanillin melts at about $80^{\circ}C$ ($176^{\circ}F$), the *U S P* states between 80° and $81^{\circ}C$ (176° and $177^{\circ}F$) and that at $253^{\circ}C$ ($546^{\circ}F$) it can be

distilled without decomposition in a current of Carbon Dioxide. It is sparingly soluble in Water, but dissolves readily in Alcohol, Ether and Chloroform, also in aqueous solutions of alkali Hydroxide, from which latter solution it is reprecipitated on neutralisation of the alkali Hydroxide. The aqueous solution affords with Ferric Chloride T.S. a blue colour, changing to brown when the liquid is boiled, and affording a white precipitate on the addition of Lead Acetate Solution. This precipitate is soluble in hot Water and crystallises out in scales as the solution cools. When Vanillin is warmed with concentrated Alcohol or Sodium Hydroxide Solution, a few drops of Chloroform added, and the liquid again warmed, no odour of Phenol Isocyanide should be evolved, indicating the absence of Acetanilide.

TINCTURA VANILLÆ—Mix 65 of Alcohol (95 p.c.) with 35 of Water. Macerate 10 of Vanilla, cut small and bruised, in 50 of the mixture for 12 hours. Drain off the liquid and set it aside. Transfer the Vanilla to a mortar, beat it with 20 of Sugar into a uniform powder, then pick it in a percolator, and continue the percolation with more of the menstruum to make 100—U.S.P.

This has been incorporated in the B.P.C.

Not Official

VERATRI VIRIDIS RHIZOMA

GREEN HELLEBORE RHIZOME

The Rhizome and Rootlets of *Veratrum viride*, Aiton.

Collected in autumn in U.S. and Canada.

The principal alkaloidal constituent (about half) is **Cevadine**, the same base as is found in **Cevadilla**, **Jeivine** and **Pseudo-jeivine**, in about equal proportions, constituting the remainder.—P.J. (3) 1896.

Medicinal Properties—Sedative. Has been given to quiet spinal spasms, should be prescribed cautiously.

10 minims of the tincture with 5 grains of Chloral Hydrate given hourly, or 10 minims hypodermically, in puerperal eclampsia.—L. '98, 1 146, '99, 1 1430.

Foreign Pharmacopœias—Official in Belg., Ger., Swed. and Swiss (Rhizoma Veratri (Veratrum Album)), Mex. (Eleboro Blanco and Eleboro Verde), U.S. (Veratrum Album or Viride).

TINCTURA VERATRI VIRIDIS (L.P. '85)—Green Hellebore Rhizome, in No. 40 powder, 1, Rectified Spirit (Alcohol 88.76 p.c.) q.s. to yield 5 (1 in 5).

Dose—5 to 20 minims = 0.3 to 1.2 c.c.

The best menstruum is stated to be Alcohol (70 p.c.)—C.D. '92, 11 651.

Official in Ger., 1 in 10, U.S., 1 in 10, B.P.C., 1 in 10. U.S. has also a **Fluid Extract**, 1 in 1.

Tests—Tincture of Green Hellebore (B.P. '85) has a sp. gr. of about 0.952. It contains about 2.0 p.c. w/v of total solids and about 32.0 p.c. w/v of Absolute Alcohol.

VERATRINA.

VERATRINE

A white, or greyish-white, odourless, amorphous powder, possessing a very bitter acid taste and leaving a feeling of numbness on the tongue. It is intensely irritating to the nasal mucous membrane and the smallest particle produces violent sneezing. Permanent in the air. It is officially described as an alkaloid, or mixture of alkaloids, prepared from Cevadilla, the dried ripe Seeds of *Schœnocaulon*.

officinale, A Gray, the *USP* describes Veratrine as a mixture of alkaloids obtained from the Seed of *Asagæu officinalis*, Lindley

It should be kept in well-stoppered glass bottles of a dark amber tint and protected as far as possible from contact with the light

Commercial Veratrine is liable to be very variable in physiological activity

The nomenclature of the alkaloids contained in this mixture has undergone modification Wright and Luff assign to the crystallisable portion (called by Merck 'Veratrine') the name of **Cevadine**, as it yields on saponification Cevadic Acid, the name **Veratrine** being reserved for the base described by Couerbe, which yields Veratric Acid Another base has been called **Cevadiline**, but the bulk of the alkaloid refuses to yield any crystallisable or otherwise definable salts

Solubility.—Scarcely soluble in cold Water, 1 in 1000 of boiling Water, 1 in 3 of Alcohol (90 p c), 1 in 6 of Ether, 1 in 3 of Chloroform, sparingly in Glycerin, about 1 in 80 of Olive Oil, and readily in diluted Acids

Medicinal Properties.—A powerful irritant poison, scarcely ever given internally Externally it acts as an analgesic in ' ' ' ' , more particularly of the fifth nerve It should not be used where the skin is broken

Ph Ger maximum single dose, 0 005 gramme, maximum daily dose, 0 015 gramme

Official Preparation—Unguentum Veratrinæ

Not Official—Oleatum Veratrinæ

Antidotes—Emetic, stimulants, Coffee, warmth to the extremities Recumbent position to be strictly maintained—*Murrell*

Foreign Pharmacopœias—Official in all the Foreign Pharmacopœias, except Dan Dutch, Cevadinum

Tests.—Veratrine, *BP*, melts when heated to a yellow liquid, Veratrine, *USP*, softens at 145° C (293° F) and melts at 152° C (305 6° F), no mp is assigned to Veratrine, *PG* It dissolves in Nitric Acid, forming a yellow solution When warmed with Hydrochloric Acid it dissolves, yielding a blood-red colour permanent for some days Triturated with Sulphuric Acid it yields first a yellow and then a bright red mixture, subsequently exhibiting a yellowish-green fluorescence when viewed by reflected light, the fluorescence becoming more intense on further addition of acid If the Sulphuric Acid mixture be warmed a violet-red coloration is produced, or if it be allowed to stand a violet-red coloration is gradually produced A drop of Syrup added to the mixture of Sulphuric Acid and Veratrine darkens the red colour and gives it a purple coloration, by exposure to air the purple becomes blue Sulphuric Acid with one-seventh of its volume of Water is a more useful reagent Veratrine, *USP*, yields with Sulphuric Acid containing a trace of Selenous Acid a brick-red coloration Veratrine dissolves readily in Alcohol (90 p c), the alcoholic solution being alkaline in reaction towards Litmus paper The ' ' ' ' solution should not yield a precipitate on the addition of Platinium Chloride Solution, indicating the absence of other alkaloids such as Brucine, Morphine and Stachnine 0 5 of

a gramme when heated with free access of air should leave no weighable residue. A distinguishing reaction for Veratrine is its irritating effect upon the nasal mucous membrane, a tiny particle of the dust from the powdered alkaloid causing violent sneezing. The test should, however, be applied with extreme caution, and the same caution should be exercised in tasting substances or liquids presumed to contain the alkaloid.

UNGUENTUM VERATRINÆ VERATRINE OINTMENT

Dissolve 10 grains of Veratrine in 40 grains of Oleic Acid, at a gentle heat, and add 450 grains of Lard (1 in 50)

Now 1 in 50 instead of 1 in 63, Hard and Soft Paraffins and Olive Oil replaced by Oleic Acid and Lard

Foreign Pharmacopœias—Official in U S, 1 in 25, Port and Russ, 1 in 50

Not Official

OLEATUM VERATRINÆ (U S)—Veratrine 2, Oleic Acid 50, Olive Oil, *q s* to make 100, by weight

This has been incorporated in the *B P C* under the title **Oleatum Veratrinæ** *Syn* Oleatum Veratrinæ

Squibb suggests that this should be made 10 p c as more likely to give relief in neuralgia.—*Squibb*, p 164

Not Official

VIBURNUM

BLACK HAW

The Bark of *Viburnum prunifolium*, L

It is official in the *Ind* and *Col Add* for India and the Eastern and North American Colonies, also **Extractum Viburni Prunifolii Liquidum** (1 in 1) Dose, 60 to 120 minims = 3 6 to 7 1 c c

Medicinal Properties—Strongly recommended as a preventive in cases of threatened abortion, to control menorrhagia and metrorrhagia and in all kinds of pelvic inflammation, brilliant results in dysmenorrhœa.—*M* 4 '95, 192, *B M J* '95, ii 1562, *L* '95, ii 1625

Foreign Pharmacopœias—Official in Austri, Dutch, Fr, Mex, Span and U S

The bark of *Viburnum opulus* has also been used in similar cases

ELIXIR VIBURNI PRUNIFOLII—Fluid Extract of *Viburnum Prunifolium*, 12 5, Compound Tincture of Cardamom, 7 5, Aromatic Elixir, 80 Average dose, 1 fl drim = 3 6 c c—*U S N F*

This has been incorporated in the *B P C*

ELIXIR VIBURNI PRUNIFOLII COMPOSITUM—Liquid Extract of *Viburnum Prunifolium*, 50, Dry Extract of Hydrastis, 1 75, Oil of Coriander 0 50, Oil of Calaway, 0 50, Glycerin, *q s* to produce 100—*B P C*

EXTRACTUM VIBURNI PRUNIFOLII LIQUIDUM—Percolate 20 of Black Haw, in No 60 powder, with Alcohol (70 p c) until exhausted, reserve the first 17, reduce the remainder to a soft extract, dissolve this in the reserved portion, and add Alcohol (70 p c) *q s* to make 20—*Ind* and *Col Add*

Dose—60 to 120 minims = 3 6 to 7 1 c c

This has been incorporated in the *B P C*

This Fluid Extract evaporated to a firm extract constitutes **Extractum Viburni Prunifolii**—*B P C*

FLUIDEXTRACTUM VIBURNI PRUNIFOLII (*U S*) --Exhaust by percolation Viburnum, in No 40 powder, 100 parts, with a mixture of Alcohol (95 p c), 2, and Water, 1, reserve the first 85, and evaporate the remainder to a soft extract, dissolve this in the reserved portion, and add enough menstruum to measure 100

Foreign Pharmacopœias—Official in Austr, Dutch and Fl, 1 in 1
U S has also Fluidextractum Viburni Opuli, 1 in 1

VINA.

WINES

Medicated wines are of very ancient date, and were admitted to our earliest Pharmacopœias. Two only remain as representatives of the old Pharmacopœias—Vinum Antimoniale and Vinum Ferri, the former was prepared by digesting 4 oz of the Regulus of Antimony in powder with 3 lb of 'White' Wine (Pharmacopœia Londinensis, 1655). The latter (Vinum Chalybeatum) was made with Rhensish Wine and Iron filings

VINUM XERICUM.

SHERRY

A Spanish Wine

Unless good sound Sherry is used, the preparations are apt to spoil by keeping

It contains about 20 p c Alcohol by volume

Official Γ Used in the preparation of Vinum Antimoniale,
Vinum Colchici and Vinum Ipecacuanhæ

Not Official—Vinum Xericum Detannatum

Tests—Sherry of good quality has a sp gr of about 0.985 to 0.998. It is officially required to contain not less than 16 p c by volume of Ethyl Hydroxide. Good sound Sherries contain from 16 to 20 p c by volume of Absolute Alcohol, the Alcohol may be determined by a similar method to that given under Spiritus Frumenti. The total acid usually amounts to about 0.52 p c w/v calculated as Tartaric Acid, that is to say, a measured quantity of 10 c c of the Wine will require about 7.0 c c of Decinormal Volumetric Sodium Hydroxide Solution for neutralisation, Phenolphthalein Solution being used as an indicator of neutrality. The extractive matter may vary from 2 to 5 p c w/v. The ash amounts to about 0.55 p c w/v. The Wine is officially required to be free from Salicylic Acid. The official method of testing being as follows—A measured quantity of 50 c c is mixed with 50 c c of Water, 5 c c of Normal Volumetric Sulphuric Acid Solution added, and the mixture distilled. The first 10 c c portion of the distillate is rejected, the balance is shaken with Ether, the ethereal liquid separated and the Ether removed by evaporation. The residue is required to yield no violet coloration on the addition of Ferric Chloride T.S. Theoretically considered the test appears unsatisfactory. The first 10 c c portion may possibly contain Ethyl Salicylate passing

over with the spirit, the evaporation of the ethereal Solution of the Salicylic Acid is not to be recommended owing to the risk of loss by evaporation. A preferable plan would have been to have added sufficient Water to the ethereal liquid to form a separate layer, and 1 or 2 drops of Ferric Chloride T.S. and to shake vigorously, if Salicylic Acid be present the lower aqueous layer is coloured an immediate violet. A useful test for the presence of Salicylic Acid is given under Vinum Auranti.

Not Official.

VINUM XERICUM DETANNATUM (*B.P.C.*)—Sherry, 100, Gelatin, in No. 100 powder, 0.15, macerate for 24 hours (at a temperature not exceeding 15° C.) with frequent agitation, and decant.

Bud has shown (*Y.L.P.* '99, 363) that by substituting Gelatin in No. 100 powder (now commercially procurable) for Gelatin cut small, as previously directed in the *B.P.C. Formulary* 1894, it is possible to completely detannate an average sample of Sherry in 24 hours. The same Wine treated with sheet Gelatin cut small required days for the completion of the process.

Not Official

VINCA MAJOR

GRIATRE PERIWINKLE

An infusion made of dried Herb 2 boiling Water 20, is powerfully astringent, and will often arrest menorrhagia.

Dose—A wineglassful.

Foreign Pharmacopœias—Official in Fr. (*Peirvenche Officinale*).

Dose of the fluid extract, 1 to 2 fl. dm. = 3.6 to 7.1 cc.

Not Official

VIOLA

The flowers of *Viola odorata*, L., are official in the French, Portuguese and Spanish Pharmacopœias.

The herb *Viola tricolor*, L., is official in the Austrian, German and Swiss Pharmacopœias. That official in the Austrian is the cultivated variety, that in the German and Swiss from wild plants.

A certain amount of interest is attached to the leaves of the Violet on account of an apparent improvement following the employment of the fresh infusion of the leaves in a case (*L.* '05, 1 713) in which it was alleged that a patient might have been suffering from malignant disease. A handful of the leaves was soaked in a pint of boiling Water for 24 hours and the liquid poured off, divided into 2 parts, 1 part being taken internally during the 24 hours and the other used as a fomentation. An apparent recovery from a presumably malignant growth of the mouth resulted. An examination of the leaves of the common Violet (*Viola odorata*) in the *Lancet* laboratory (*L.* '05, 1 1085) showed the presence of two crystalline bodies, one glucosidal and the other alkaloidal in character, and also a dark green Oil. Alcohol was found a much more effective solvent than an aqueous menstruum, in view of the employment of an aqueous infusion the latter point is of interest. The alkaloid isolated behaved chemically much in the same way as Emetine, the principal alkaloid of *Ipecacuanha*. It has been stated (*Y.L.P.* '05, 467, *C.D.* '05, 11 977, *P.J.* '05, 11 869) that any activity which Violet leaves possess is due either to the glucoside, the product of its decomposition, or a natural ferment associated with it. Reckoned as *Viola quercitina*, the glucoside from Princess of Wales Violet leaves amounted to 5 p.c. of the weight of the fresh leaves. A fresh infusion was found to extract nine

tenths of the glucoside present in the leaves. No volatile constituent was isolated, no alkaloid could be detected, no Salicylic Acid was found. The presence of a glucoside was proved, but the glucoside was not isolated. Objection has been taken to the evidence of the uses of Violet leaves having been unfortunately collected chiefly by unskilled persons, and that it has therefore been lacking in definiteness, and consequently in value. After the definite expression of the opinions mentioned in the above reference, it is to find in a paper read before the Therapeutical Society, October 2, 1896, and reported in the *Lancet*, '06, ii 1318, that 'isolato and identify a glucoside from Violet leaves have failed, was no evidence of a ferment being present, the only positive fact resulting from the experiments being that the leaves and their preparations yield under certain conditions glucose.'

It has been pointed out that the reputation of Violets for the treatment of malignant growths was founded on the use of wild Violets, at least as far back as James I., and that it is therefore desirable that in any inquiry into the subject wild Violets should be used, such as have been used for centuries, and not a recent cultivated Violet as employed at the present time. In the light of the above remarks, the varieties official in the Continental Pharmacopœias will be of interest. It will be noted that wild Violets are official in the German and Swiss Pharmacopœias, and cultivated Violets in the Austrian.

Not Official

YEAST

BEER YEAST

The ferment obtained in brewing Beer and produced by *Saccharomyces cerevisia*

A viscid, frothy semi-fluid, possessing a sour vinous odour and a somewhat bitter taste. It is insoluble in Alcohol, practically insoluble in Water. Exposed to a moderate heat it loses its liquid portion and becomes dry, hard and brittle, and in this form may be preserved for some time, though apparently with a loss of much of its peculiar power. Yeast cakes are prepared by putting Yeast into sacks, washing with Water, submitting it to pressure, and ultimately drying it. Compressed Yeast, the undried product, is now largely used.

Medicinal Properties—Antiseptic and stimulating, it has been recommended internally as a proteolytic against boils and carbuncles, and has been found useful in obstinate dysentery. In typhoid fever (*L* '05, i 463) 60 grammes daily, in 3 doses, commenced about the seventh day, to improve the gastro-intestinal symptoms, to reduce the temperature and diminish diarrhoea. Living Yeast does not possess any directly bactericidal or phagocytic properties. Injected intravenously it causes intravascular clotting of the blood. Subcutaneous injections of pure cultures of living Yeast can be made in animals without producing any ill effects. Killed Yeast produces the same effects as living Yeast. The immediate effect of subcutaneous injections is to produce leucopenia, rapidly followed by the leucocytosis. The effects produced by the injections of Yeast are probably due to the nucleo albumen contained in the cells of the body generally, and cause a large increase in the antiseptic and anti-bactericidal substances

the blood serum—*B M J Supplement* '05, ii 7
and acne—*F T* '07, 19

Dose— $\frac{1}{2}$ to 1 oz. alone or with Water

Furunculine and Levurine are powdered forms of dehydrated Yeast

Not Official

YOHIMBINE.

Silky white needles, or as a white inodorous amorphous powder, which has a tendency to change in colour on exposure to light, it should therefore be kept in well-closed glass bottles of a dark amber tint and protected as far as possible from

the light. It possesses a faint odour of Benzaldehyde. It is slightly soluble in Water, readily soluble in Methyl, Ethyl, or in Amyl Alcohol, in Ether and in Chloroform. It is an alkaloid derived from the bark of *Corynanthe yohimbe* (Schumann) or Yohimbe tree, which grows in the Southern Cameroons district in Africa.

It is said to act as an aphrodisiac. It causes anesthesia of the cornea and conjunctiva when dropped into the conjunctival sac. It is stated (*L* '05, 1 1013) to be useful in chronic affections of the eye which require stimulation. It is preferred to Tiopacocaine, as its effects are more persistent, and to Cocaine, as it does not affect the epithelium, interfere with the nutrition of the cornea or produce mydriasis and hypotony, and is, moreover, non-toxic. 25 cc of a 1 p c solution may be injected subcutaneously without harm, producing a local anesthesia which lasts for nearly 2 hours. It has been found (*L M J E* '05, 1 28) useful in cases of toxic impotence. It is reported to have a favourable influence in cases of neurasthenic impotence.

Tests—Yohimbine melts at about 234° C (453 2° F). According to Arnold and Behrens (*Pharmazeutische Zentralblatt*, xlii 49) it has certain properties in common with Cocaine. It produces a temporary anesthesia somewhat resembling that occasioned by Cocaine. They give the following reactions for distinguishing between the two.—The m p, Cocaine melts at 98° C (208 4° F), Yohimbine as stated above, Cocaine Hydrochloride melts at 183° C (361 4° F), Yohimbine Hydrochloride has a m p as given below, Cocaine when heated for 5 minutes with Sulphuric Acid yields an odour of Methyl Benzoate, Yohimbine yields a faint odour resembling Peppermint, Cocaine when heated first with Fuming Nitric Acid and then with Hydrochloric Acid Solution gives no colour reaction, Yohimbine is coloured at first a deep green and then yellow by Nitric Acid, on the addition of Alcoholic Potassium Hydroxide Solution a cherry red colour is produced, Cocaine remains colourless when dissolved in strong Sulphuric Acid, and when treated with Chlorinated Lime, Yohimbine gives an intense orange red colour, Cocaine gives a black coloration when triturated with Mercuric Chloride, Yohimbine produces no such black coloration. When dissolved in strong concentrated Sulphuric Acid it affords on the addition of a minute crystal of Potassium Bichromate a beautiful violet coloration. It yields with Cane Sugar and Sulphuric Acid a wine red colour. Attention, however, has been called to the fact that Saccharose, Glucose or Furfural by themselves afford, with Sulphuric Acid, a red or reddish violet coloration, and that Sesame Oil also produces a similar reaction. This colour reaction, therefore, cannot be regarded as serviceable for the identification of Yohimbine. 0.1 of a gramme when ignited with free access of air should leave no weighable residue.

YOHIMBINE HYDROCHLORIDE—It occurs in colourless crystals, slightly soluble in Water. It is the Hydrochloride of the alkaloid Yohimbine.

It should be kept in well closed bottles of a dark amber tint and protected as far as possible from contact with the light.

Tests—Yohimbine Hydrochloride melts at 290° C (554° F), Cocaine Hydrochloride melts at 183° C (361 4° F). An aqueous solution affords an amorphous greyish violet precipitate on the addition of Auric Chloride Solution (1 p c), an aqueous solution of Cocaine Hydrochloride yielding on the addition of the same reagent a pale yellow precipitate of microscopic needles.

Not Official YERBA SANTA

The dried Leaves of *Eriodictyon Californicum* (Hook and Arn.), Greene, are official in the *U S P*. They contain about 30 p c of resin, some essential oil, Glucose, two hydrocarbons, fatty acids, Phytosterol and three crystalline substances of a phenolic nature.—*J C S Abs* '06, ii 885.

A stimulating expectorant, recommended in acute bronchitis.

Fluidextractum Eriodictyi (U S), 1 in 1, with a mixture of Alcohol (95 p c) 4 and Water 1, average dose, 17 minims = 1 cc.

Not Official
Z I N C U M

ZINC

Zn, eq 64 91

A bluish-white metal, of peculiar taste and of a perceptible smell when rubbed, laminated, and with a crystalline fracture

It occurs native, as a Sulphide or as a Carbonate, and is separated from impurities by sublimation

The laminated or granulated metal is official in the Appendix to the *BP* Metallic Zinc is official in the body of the *USP*. It appears in the lists of Reagents in the Appendix to the *PG*. The *USP* states that it is in the form of thin sheets, in irregular granulated pieces, or moulded into thin pencils or in fine powder

Official Preparations—Used to prepare Liquor Zinci Chloridi, Zinci Chloridum, Zinci Oxidum, Zinci Sulphas

Foreign Pharmacopœias—Official in Mex, Span and U S

Tests—Zinc has a sp gr of 7.1, the *USP* states from 6.9 when it is cast to 7.2 when rolled. It fuses at 411.6° C (773° F). The *USP* states 412° to 779° F. At about 940° C (1724° F) it boils, and may be readily distilled. It dissolves readily in diluted Hydrochloric Acid, simultaneously evolving Hydrogen gas, which burns with a blue flame on ignition. It yields, when dissolved in Hydrochloric Acid, a clear solution, which should answer the following tests.—When neutralised with Ammonia Solution it yields with Ammonium Hydrosulphide Solution a white precipitate insoluble in Acetic Acid, soluble in Hydrochloric Acid. When mixed with sufficient Ammonium Chloride to hold in solution the Hydroxide which would otherwise be precipitated, it yields on the addition of Ammonia Solution a white precipitate, it affords with Hydrogen Sulphide a similar white precipitate, insoluble in Acetic Acid, soluble in Hydrochloric Acid, with Ammonia Solution, Potassium or Sodium Hydroxide Solution it affords a white precipitate soluble in excess of the reagent, with Potassium Ferrocyanide Solution it yields a white precipitate insoluble in diluted Hydrochloric Acid. It should not contain Antimony, Arsenic, Aluminium, Cadmium, Copper, Iron, Lead or Magnesium, Sulphur or Phosphorus. A measured quantity of 10 c.c. of a solution containing 5 p.c. of the metal in diluted Hydrochloric Acid (a slight excess of Hydrochloric Acid being present) should not afford a coloration or turbidity when mixed with an equal volume of freshly prepared saturated Hydrogen Sulphide Solution and allowed to stand for 30 minutes, indicating the absence of Antimony, Arsenic, Cadmium, Copper and Lead. The solution in dilute Hydrochloric Acid when mixed with Ammonium Chloride and Ammonia Solution and boiled should not yield either a flocculent white precipitate or a brown flocculent precipitate, indicating the absence of Aluminium and Iron. A further portion of a similar solution mixed with Ammonium Chloride and Ammonia Solution should not afford a turbidity on the addition of either Ammonium Oxalate Solution or Sodium Phosphate Solution, indicating the absence of Calcium and Magnesium. The specimen should not yield a reaction for Arsenic when examined by the modified Gutzeit's test or by the Bettendorf's test. The Hydrogen gas evolved during the solution of the metal in diluted Hydrochloric Acid should not possess the distinctive disagreeable odour of Hydrogen Sulphide, nor should a strip of Lead Acetate paper be altered in colour when suspended in the issuing gas, indicating the absence of Sulphur. A strip of paper moistened with Silver Nitrate Solution when similarly held in the escaping gas should not be blackened, indicating the absence of Phosphorus, and affording confirmatory evidence of the absence of Antimony, Arsenic and Sulphur. In performing the time-limit test for Arsenic, Cadmium, Copper, Lead and Iron, the *USP* dissolves 1 gramme of Zinc in a mixture of 10 c.c. of Nitro-hydrochloric Acid and 10 c.c. of Water, evaporates the solution to dryness, moistens the residue with 2 c.c. of Hydrochloric Acid, again evaporates to dryness, and finally dissolves the residue in 10 c.c. of Water.

ZINCI ACETAS.

ZINC ACETATE

Zn (C₂H₃O₂)₂, 3H₂O, eq 235 71FR, ACETATE DL ZINC GLR, ZINACHLORID, ITAL, ACIATO DI ZINCO,
SPAN, ACIATO ZINCICO

Soft white glistening monoclinic crystals, possessing an acetous odour and a sharp metallic taste

Zinc Acetate official in the *U S P* contains 2 molecules of Water of crystallisation, that official in the *B P* 3 molecules. The *U S P* Acetate is required to contain in the uneffloresced condition not less than 99.5 p.c. of pure crystallised Zinc Acetate

It should be kept in well closed vessels, as it has a tendency to effloresce on exposure to air, and also to lose Acetic Acid with the formation of a basic salt

Solubility—10 in 25 of Water, 4 in 1 of boiling Water, 1 in 40 of Alcohol (90 p.c.), 1 in 3 of boiling Alcohol (90 p.c.)

Medicinal Properties—Similar to the Sulphate, chiefly used as a local astringent

Dose—1 to 2 grains = 0.06 to 0.13 gramme

Not Official—Lotio Zinci Acetatis

Foreign Pharmacopœias—Official in Ger, Hung, Mex, Port, Russ and U S

Tests—Zinc Acetate when heated partially fuses, losing its Water of crystallisation and a certain amount of acid. At still higher temperatures it is decomposed, and when ignited at a dull red heat leaves a residue of Zinc Oxide. It dissolves readily in Water, forming a solution which is slightly acid in reaction towards blue Litmus paper, but which is not always clear, as the commercial salt sometimes contains a small proportion of basic salt. It affords, however, a clear solution on the addition of a little Acetic Acid. The solution answers the tests distinctive of Zinc given under that heading. The aqueous solution affords on the addition of Ferric Chloride Solution a red coloration, changing to a reddish brown precipitate on boiling. When warmed with Sulphuric Acid it evolves a distinctive acetous odour. When warmed with Sulphuric Acid containing a little Alcohol (90 p.c.) it evolves the peculiar odour of Ethyl Acetate (Acetic Ether). The dry salt, heated with a minute proportion of Arsenious Anhydride, yields the distinctive but highly poisonous odour of Cacodyl Oxide. The impurities mentioned under Zinc should be absent. Solutions should answer the tests given under the headings of Barium Chloride, Silver Nitrate, Sulphuric Acid, and Gutzzeit's test appearing in small type below, indicating the absence of Sulphates, Chlorides, impurities derived from the use of Acetic Acid containing empyreumatic impurities and Arsenic

Hydrogen Sulphide—In an aqueous solution (1-10) of the salt Hydrogen Sulphide T.S. produces a pure white precipitate. The liquid filtered off from the

precipitate should not leave a weighable residue on evaporation, *P G* 10 c c of an aqueous solution (1-20), to which 1 c c of Hydrochloric Acid has been added, should not respond to the time-limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted, *U S P*

Sulphuric Acid—On gently warming the salt with Sulphuric Acid it should not undergo any blackening, *P G*

Barium Chloride—An aqueous solution of the salt (1-20) after the addition of a few drops of diluted Nitric Acid should remain clear upon the addition of T S of Barium Chloride, *U S P*

Silver Nitrate—An acidulated solution as above should remain clear on the addition of T S of Silver Nitrate, *U S P*

Gutzeit's Test—5 c c of the aqueous solution of the salt (1-10) should not respond to the modified Gutzeit's test for Arsenic, *U S P*

Not Official

LOTIO ZINCI ACETATIS—Zinc Acetate, 2 grams, Water, 1 fl oz *Mix*
An astringent collyrium in conjunctivitis, or as an injection in gonorrhoea after the acute stage has passed

Tincture of Opium causes no precipitate with this Lotion

A lotion very commonly prescribed at one time was that containing Zinc Sulphate and Lead Acetate, which mutually react with formation of soluble Zinc Acetate and insoluble Lead Sulphate, it has been superseded by the above.

Not Official

ZINCI BROMIDUM.

A whitish, very deliquescent, granular powder

It should be kept in well-stoppered glass bottles and exposed as seldom as possible to the air, as it is extremely deliquescent

The *U S P* salt is required to contain when anhydrous at least 97 p c of pure Zinc Bromide

Solubility—4 in 1 of Water, 2 in 1 of Alcohol (90 p c)

Dose—2 grains = 0.13 gramme 3 times a day for epilepsy

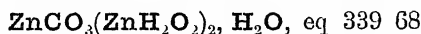
Official in Mex, Span and U S

Tests—Zinc Bromide when strongly heated fuses, the *U S P* states at a temperature of 394° C (741.2° F). It dissolves readily in Water, yielding a solution which has a slightly acid reaction towards blue Litmus paper, and which affords the tests distinctive of Zinc given under that heading. The solution also yields with Silver Nitrate Solution a yellowish curdy precipitate insoluble in Nitric Acid, when separated and washed. It is soluble in Ammonia Solution, but readily soluble in Potassium Permanganate. When heated with Sulphuric Acid and Manganese Dioxide, reddish vapours of Bromine are evolved, which produce an orange-yellow stain on filter paper moistened with Starch Mucilage. It is required by the *U S P* to contain not less than 97 p c of pure Zinc Bromide as volumetrically determined by dissolving a quantity of 0.3 of a gramme of the anhydrous salt in 10 c c of Water, a Tenth-normal Volumetric Silver Nitrate Solution, using Potassium Chromate Solution as an indicator, not less than 26 c c nor more than 26.8 c c should be required to produce the end reaction. The impurities mentioned under Zinc should be absent from Zinc Bromide. The *U S P* fixes a limit of Chloride, not more than 1 gramme dissolved in 50 c c of Acetic Acid and 2 grammes of Hydrochloric Acid (free from Chloride) added, the mixture evaporated in a small beaker to at least 10 c c, the residue diluted with 10 c c of Distilled Water and filtered, should not give more than a slight turbidity on the addition of 2 c c of Nitric Acid and a few drops of Silver Nitrate Solution. The aqueous solution of the salt when mixed with Chlorine Water, diluted with an equal volume of

Water, and shaken with Carbon Bisulphide, the latter solution should not assume a violet colour, indicating the absence of Iodide. The aqueous solution of the salt should not yield a turbidity on the addition of Barium Chloride Solution, indicating the absence of Sulphate.

ZINCI CARBONAS.

ZINC CARBONATE ZINC HYDROXYCARBONATE



FR, CARBONATE DE ZINC, GÉR, ZINKCARBONAT, ITAL, CARBONATO DI ZINCO
SPAN, CARBONATO ZINCICO

A dry, white, odourless and tasteless amorphous impalpable powder, permanent in the air.

It may be produced by precipitating solution of Zinc Sulphate with Sodium Carbonate. The precipitated Carbonate official in the *U S P* is the hydrated Zinc Carbonate, and is required to yield, on ignition, not less than 72 p c of Zinc Oxide. The Carbonate is not official in the *P G*.

The anhydrous normal Carbonate, ZnCO_3 , occurs native as Calamine. The composition of the precipitated hydrated Carbonate varies much according to the conditions under which it is formed.

Medicinal Properties—A mild astringent, used with other substances as a dusting powder, also in lotions.

Official Preparations—Used in the preparation of Zinci Acetas, Zinci Oxidum, and Zinci Valerianas.

Foreign Pharmacopœias—Official in U S, Zinci Carbonas Precipitatus.

Tests—Zinc Carbonate when strongly heated loses Water and Carbon Dioxide, leaving a residue which whilst hot is yellow, and which when cold is white. It dissolves readily and completely with effervescence in Diluted Nitric Acid, and yielding a gas which, when passed into Lime Water, affords a white precipitate soluble in a sufficient excess of the gas, or soluble with effervescence in Diluted Hydrochloric Acid. The solution in Diluted Hydrochloric Acid answers the tests distinctive of Zinc given under that heading. The *BP* does not require it to yield any definite percentage of Oxide upon ignition. The *U S P* requires it to yield not less than 72 p c, 1 gramme of the salt when strongly ignited being required to yield a residue weighing not less than 0.72 gramme. The impurities mentioned under Zinc should also be absent. A solution in Diluted Nitric Acid should not afford a pronounced turbidity with either Silver Nitrate Solution or Barium Chloride Solution, indicating the absence of more than traces of Chloride and Sulphate. The *U S P* fixes a limit of alkali calculating out to 0.3 p c of anhydrous Sodium Oxide, the test is described in small type below under the heading of Phenolphthalein. The solution employed by the *U S P* in carrying out the tests of identity and purity is obtained by mixing 1.25 grammes of the salt with 10 c c of Diluted Sulphuric Acid and 10 c c of Water, removing the undissolved excess by filtration after effervescence has ceased.

Time-limit Test—Add 10 c c of diluted Sulphuric Acid and 10 c c of Water to 1.25 grammes of the salt, and after effervescence has ceased, remove the undissolved excess by filtration. A portion of the filtrate acidulated with Hydrochloric Acid should not respond to the time-limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted. *U S P*

Phenolphthalein—If 1 gramme of the salt be placed in a flask with 10 c c of boiling Water, and 2 drops of Phenolphthalein T S added, not more than 1 c c of Tenth-normal Hydrochloric Acid Volumetric Solution should be required to discharge the red colour, *U S P*

ZINCI CHLORIDUM.

ZINC CHLORIDE

ZnCl_2 , eq 135.29

FR, CHLORURE DE ZINC, GER, ZINKCHLORID, ITAL, CLORURO DI ZINCO, SPAN, CLORURO ZINCICO

White or almost white, very deliquescent fused irregular masses, or peculiar-shaped sticks, or a white granular deliquescent powder. It is strongly caustic, and should be handled with great care.

Its official method of preparation is by the interaction of Zinc and Hydrochloric Acid.

It should be kept in small, well-stoppered glass bottles and exposed as little as possible to the air, as it is extremely deliquescent.

Solubility.—10 in 4 of Water, 1 in 1 of Alcohol (90 p c), freely in Ether, 1 in 4 (nearly) of Glycerin.

Medicinal Properties—Diluted it is antiseptic and disinfectant. Seldom given internally. Externally, applied as a caustic, in form of point or paste, to indolent ulcers and malignant growths, to condylomata, and to naevi. As a lotion, 20 grains to 1 fl oz of Water, it is an efficient substitute for Carbolic Acid, in syringing offensive pus cavities, sinuses, foul ulcers, etc.

As a paste for packing the cavity of uterus in malignant disease—*B M J* '95, 1756

As an injection (1 grain to 1 fl oz) in gonorrhoea.

Official Preparation—Liquor Zinci Chloridi

Not Official—Zinc Chloride Points, Cautique au Chlorure de Zinc, Guttæ Zinci, Zinci Chloridi cum Cocaina, Lotio Zinci Chloridi, Pasta Zinci Chloridi, Pasta Zinci Chloridi Comp, Pulvis Zinci Chloridi Comp

Antidotes—See Zinc Sulphas, p 1250

Foreign Pharmacopœias—Official in Aust, Belg, Dan, Dutch, Fl, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests.—Zinc Chloride fuses when heated. The *U S P* says at a temperature of 115° C (239° F). The *B P* states that it is almost entirely soluble in Water, Alcohol (90 p c) and Ether. The commercial salt frequently contains a small proportion of Oxysulphide, and does not then yield clear solutions with these solvents. The *U S P* states that the 1 in 20 aqueous solution should be clear, or at the most only faintly opalescent, and if mixed with an equal volume

of Alcohol (94.9 p.c.), a single drop of Hydrochloric Acid should suffice to render 10 c.c. of the mixture perfectly clear. The *P.G.* states that the 1 in 20 aqueous solution should be clear, or at the most but faintly turbid, and that the flocculent precipitate resulting on the addition of 3 volumes of Alcohol (90 p.c.) to this solution should again disappear on the addition of 1 drop of Hydrochloric Acid. The aqueous solution answers the tests distinctive of Zinc given under that heading. On the addition of Silver Nitrate T.S. it affords a white curdy precipitate, which, when separated and washed, is insoluble in Nitric Acid, readily soluble in Ammonia Solution or Potassium Cyanide Solution. When the salt is heated with Manganese Dioxide and Sulphuric Acid it evolves a greenish-yellow gas, possessing a distinctive pungent odour, and which produces a blue coloration with paper soaked in Starch Mucilage and Potassium Iodide Solution. Neither the *B.P.* nor the *P.G.* states that it should contain any definite percentage of Zinc Chloride. The *U.S.P.* requires that it shall contain not less than 99.5 p.c. of pure Zinc Chloride as gravimetrically determined by dissolving 0.5 of a gramme of the salt in 200 c.c. of boiling Water, adding 5 drops of Phenolphthalein T.S. and sufficient Sodium Carbonate T.S. with constant stirring to yield a permanent alkaline reaction, the resulting precipitate is transferred to a filter and washed with boiling Water until all soluble matter is dissolved, dissolved in a sufficient quantity of Nitric Acid, evaporated to dryness and ignited until constant in weight, the residue should weigh not less than 0.297 gramme. The impurities mentioned under Zinc should be absent from the Zinc Chloride. It should respond to the test given in small type below under the heading of Barium Chloride, indicating the absence of Sulphates, and should also answer the tests described below under the heading of Ammonia Solution and Hydrogen Sulphide, indicating the absence of Calcium, Magnesium and alkali impurities.

Hydrogen Sulphide—An aqueous solution (1-10) should not become coloured by T.S. of Hydrogen Sulphide after the addition of Hydrochloric Acid, *P.G.* An aqueous solution (1-20) with 1 c.c. of diluted Hydrochloric Acid added should not respond to the time limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted, *U.S.P.*

Barium Chloride—An aqueous solution (1-20) after the addition of 1 c.c. of diluted Hydrochloric Acid should not be rendered turbid by the addition of T.S. of Barium Chloride, *U.S.P.*

A (1-10) aqueous solution of the salt should not be rendered turbid on the addition of T.S. of Barium Nitrate, *P.G.*

Ammonia Solution and Hydrogen Sulphide—1 gramme of the salt should give a clear solution with 10 c.c. of Water and 10 c.c. of Ammonia T.S., and this solution should give a pure white precipitate with excess of Hydrogen Sulphide T.S. The liquid filtered off from this precipitate after evaporation and heating to redness should not leave a weighable residue, *P.G.*

LIQUOR ZINCI CHLORIDI SOLUTION OF ZINC CHLORIDE

16 of granulated Zinc dissolved by heating with 44 of Hydrochloric Acid mixed with 20 of Distilled Water, and evaporated to 40. It should be free from Iron and Lead.

When made as above the solution will be basic and precipitate Oxychloride on dilution with Water. It should be evaporated rather lower, then neutralised

with Hydrochloric Acid (so that it will cease to precipitate on being diluted with 10 volumes of Water, or when this diluted solution just reddens Methyl Orange Solution), and finally made up to 40

When finished without loss the above quantities will yield a solution sp gr about 1.53. For details and an improved formula of Chlor-Zinc Iodine (Schulze's Solution) see *P J* (3) xxiii 648

Official in U S, sp gr 1.548 at 25° C (77° F)

Fr Codex states that the commercial solutions vary from 1.26 to 1.81, that having a sp gr of 1.45 is to be preferred

Tests.—Solution of Zinc Chloride has a sp gr of about 1.53. The *B P* states 1.530. The *U S P* solution contains about 50 p c by weight of Zinc Chloride, and is required to possess a sp gr of about 1.548 at 25° C (77° F). It should answer the tests distinctive of Zinc given under that heading and of Chlorides given under Zinc Chloride. It should not contain the impurities mentioned under Zinc, and when diluted should yield no turbidity on the addition of Barium Chloride T S, indicating the absence of Sulphate

Not Official

ZINC CHLORIDE POINTS—Zinc Chloride fused and run into conical moulds, preserved in glass tubes

Darts of Zinc Chloride have been used in the treatment of anthrax—*B M J* '87, ii 644

COMPOUND ZINC CHLORIDE POINTS—Zinc Chloride, 1, Zinc Oxide, 1, Wheaten Flour, 2, Water to make a stiff paste, which is formed into points

CAUSTIQUE AU CHLORURE DE ZINC—Zinc Chloride, 32, Zinc Oxide, 8, Dried Wheaten Flour, 24, Distilled Water, 4—*Fr*

GUTTÆ ZINCI CHLORIDI—Zinc Chloride, 2 grains, Distilled Water, 1 fl oz—*London Ophthalmic*

GUTTÆ ZINCI CHLORIDI CUM COCAINA—Zinc Chloride, 2 grains, Cocaine Hydrochloride, 10 grains, Distilled Water, 1 fl oz—*London Ophthalmic*

LOTIO ZINCI CHLORIDI—Zinc Chloride, 1 grain, Distilled Water, 1 fl oz—*London Ophthalmic*

This has been incorporated in the *B P C* as follows—Zinc Chloride, 1, Distilled Water, 400

PASTA ZINCI CHLORIDI—Zinc Chloride, 1, 2 or 4, Starch, 6, Lard, 1, Glycerin of Starch, *q s*, rub the Zinc Chloride with the Lard and Starch, and make into a thick paste with Glycerin of Starch—*University*

PASTA ZINCI CHLORIDI COMPOSITA—Zinc Chloride paste, 94, Extract of Opium, 4, rub the Extract smooth with 2 of Water, and then mix thoroughly with the paste—*University*

PULVIS ZINCI CHLORIDI COMP—Zinc Oxide, mixed with an equal weight of Zinc Chloride, will preserve the latter dry enough to blow through a tube into any cavity required, and may be so kept in a bottle for a long time

Not Official

ZINC IODIDUM.

A white or whitish powder, which rapidly becomes moist and changes to a brown colour on exposure to the air. On account of its nature and its liability to thus change, it should be kept in a bottle of a dark amber tint and protected as far as possible from the light. It is readily soluble in Water, Glycerin, or Ether.

It has been used as an alterative remedy in chorea, scrofula and hysteria, but has not come into general use. It has also been employed as an external application.

Tests—Zinc Iodide dissolves readily in Water, forming a solution which is acid in reaction towards blue Litmus paper, this solution should answer the tests distinctive of Zinc given under that heading. An aqueous solution yields with Silver Nitrate Solution a curdy, yellow precipitate insoluble in Nitric Acid, practically insoluble in Ammonia Solution, but soluble in Potassium Cyanide Solution, with Mercuric Chloride Solution it yields a brilliant scarlet precipitate soluble in Potassium Iodide Solution. The salt is official in the *U S P*, which requires it to contain, when anhydrous, not less than 98 p c of pure Zinc Iodide as volumetrically determined by dissolving a weighed quantity of 0.5 of a gramme of the dry Iodide in 20 c c of Water, adding 35 c c of Tenth normal Volumetric Silver Nitrate Solution, 5 c c of Nitric Acid and 3 c c of Ferric Ammonium Sulphate T S, shaking the mixture well and titrating the excess of Tenth normal Volumetric Silver Nitrate Solution with Tenth normal Volumetric Potassium Sulphocyanate Solution, not less than 3.4 nor more than 4 c c should be required.

Dose— $\frac{1}{2}$ to 2 grains = 0.032 to 0.13 gramme

Foreign Pharmacopœias—Official in Mex and U S, not in the others

Not Official

ZINCI NITRAS

Large, colourless, deliquescent, prismatic crystals, very soluble in Water and in Alcohol (90 p c)

Medicinal Properties—Used as a caustic in the place of Zinc Chloride, it penetrates deeper and produces less pain.

It can be made into a paste in the same way as Zinc Chloride.

Tests—Zinc Nitrate answers the tests distinctive of Zinc given under that heading, and should also be free from the impurities usually occurring in the metal and referred to in the text. When Ferric Sulphate Solution is carefully poured upon the surface of a cooled mixture of concentrated Sulphuric Acid and a solution of the salt, at the point of contact of the two liquids a dark brown zone appears.

When ignited it leaves a residue of Zinc Oxide.

ZINCI OXIDUM.

ZINC OXIDE

ZnO, eq 80.79

FR, OXYDE DE ZINC, GER, ZINKOXYD, ITAL, OSSIDO DI ZINCO,
SPAN, OXIDO ZINCICO

An odourless and tasteless, white amorphous impalpable powder, which gradually absorbs Carbonic Anhydride from the air. The method of preparation has some effect upon the colour of the product. A sample prepared from the precipitated Carbonate by ignition has a tendency to a faint yellow colour, whilst a sample prepared by the combustion of metallic Zinc is pure white.

It should be kept in well-closed vessels as it gradually absorbs Carbonic Anhydride from the air. The official process of preparation is

by the ignition of Zinc Carbonate at a dull red heat, or by combustion with metallic heat

Medicinal Properties.—Internally, but with doubtful success, as a sedative in chronic nervous spasmodic affections, and to check the perspirations of phthisis. Externally, as a mild astringent application in eczema and slight excoriations and ulcerations, in the form of ointment or paste absorbent as a dusting powder when mixed with Starch

Dose—3 to 10 grains = 0.2 to 0.65 gramme

Prescribing Notes—Generally prescribed in the form of pills. A good pill may be made by adding 'Diluted Glucose,' q.s. It is also given in lotions, with and without an equal quantity of Prepared Calamine, q.v.p. 280

Official Preparation—Unguentum Zinci. Used in the preparation of Zinc Sulphocarbolates

Not Official—Dusting Powder, Emplastrum Zinci Oxidi, Gelatinum Zinci Durum, Gelatinum Zinci, Glycér. d. Oxide de Zinc. Tassar's Paste, Pasta Unna, Pasta Zinci et Ichthamolis, Pessus Zinci, Pulvis Zinci et Bismuthi, Pulvis Zinci Oxidi Compositus, Pulvis Zinci Oxidi et Acidi Salicylici, Pulvis Zinci Oxidi et Acidi Borici, Pulvis Zinci Oxidi et Anilini, Pulvis Zinci Oleatis Compositus, Unguentum Zinci Stearatis, Unguentum Zinci cum Acido Salicylico, Pulvis Zinci et Calomelanos, Zinci Oleas (Shoomaker's), Zinc Oxide Plaster Mulls, Zinc and Salicylic Plaster Mull, and Zinc Gelatin

Foreign Pharmacopœias—Official in all, Dan., Ger. and Swiss have also Crude

Tests—Zinc Oxide when heated assumes a yellow colour which disappears on cooling. It dissolves readily and completely and with effervescence in diluted acids. When dissolved in Diluted Hydrochloric Acid the solution should answer the tests distinctive of Zinc given under that heading. The *BP* states that it should be entirely soluble when rubbed, and if necessary warmed, with Ammonia Solution mixed with strong Ammonia Solution. The 17th Edition of the *Companion* contains the following note—'It is questionable whether any commercial Zinc Oxide is entirely soluble in Ammonia.' The *BPC* states that the Oxide is never completely soluble in Ammonia Solution. Samples obtained in 1908 from the leading manufacturers were found, with one exception, to be readily and completely soluble in Ammonia. The *USP* states that it should be completely soluble in Ammonia Water. The *PG* makes no reference to its solubility in Ammonia. Neither the *BP* nor the *PG* states a requisite percentage of Zinc Oxide, nor is a method of determination indicated. The *USP* requires that it shall contain not less than 99 p.c. of pure Zinc Oxide as determined by digesting a weighed quantity of 1 gramme of the freshly ignited Zinc Oxide with 30 c.c. of Normal Volumetric Hydrochloric Acid Solution until solution is complete, adding 2 drops of Methyl Orange, and titrating with an excess of Normal Volumetric Acid Solution. Potassium Hydroxide Solution, the latter being added slowly with stirring the precipitated Oxide to redissolve before adding more acid. Not more than 5.5 c.c. of acid should be in excess. The number of c.c. of Normal Volumetric Potassium

Hydroxide Solution used is subtracted from 30, the difference represents the number of c c of Normal Volumetric Hydrochloric Acid Solution utilised in neutralising the Oxide. 1 c c of Normal Volumetric Hydrochloric Acid Solution corresponds to 4.04 p c of Zinc Oxide. The impurities mentioned under the heading of Zinc should be absent from the Oxide. It should answer the tests given in small type below under the headings of Ammonium Oxalate, Sodium Phosphate, Barium Nitrate and Silver Nitrate, indicating the absence of Calcium, Magnesium, Sulphates and Chlorides. The *USP* includes a test for limit of alkali which is described in small type below under the heading of Phenolphthalein, the figure given corresponds to 0.3 p c of anhydrous Sodium Oxide. The solubility in Ammonia referred to in the large type above is officially adopted as a test for the absence of metallic Zinc. The *USP* adopts the time limit test for the detection of Arsenic, the *PG* employs the Bettendorff's test. Standards have been suggested (*CD '08*, 1797) of 0.2 p c for Lead and 10 parts per million for Arsenic.

Ammonium Oxalate—The clear, colourless liquid obtained by dissolving 1 part of Zinc Oxide in 10 parts by weight of Diluted Acetic Acid (*PG*), when supersaturated with Ammonia Solution should not be rendered turbid by T S of Ammonium Oxalate, *PG*.

Sodium Phosphate—Neither should this liquid be rendered turbid by T S of Sodium Phosphate, *PG*.

Hydrogen Sulphide—The same liquid with Hydrogen Sulphide poured on as a layer gives a pure white zone, *PG*. Digest 1 gramme of Zinc Oxide with occasional agitation in a mixture of 10 c c of diluted Hydrochloric Acid and 10 c c of Water until saturated, then remove the undissolved Zinc Oxide by filtration. A portion of the filtrate acidulated with Hydrochloric Acid, should not respond to the time limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the Ammonia Water should be omitted, *USP*.

Phenolphthalein—If 1 gramme of Zinc Oxide be placed in a flask with 10 c c of boiling Water, and 2 drops of Phenolphthalein T S be added, not more than 1 c c of Tenth-normal Hydrochloric Acid Volumetric Solution should be required to discharge the red colour, *USP*.

Barium Nitrate—Let 2 grammes of Zinc Oxide be agitated with 20 c c of Water and the mixture filtered. The filtrate should not be rendered more than opalescent by T S of Barium Nitrate, *PG*. A solution of 1 gramme in a sufficient quantity of diluted Nitric Acid should not become more than slightly turbid upon the addition of T S of Barium Chloride, *USP*.

Silver Nitrate—The filtrate obtained as in the last test should not be rendered more than opalescent by T S of Silver Nitrate, *PG*. A solution obtained as in the last test should remain clear upon the addition of T S of Silver Nitrate, *USP*.

Stannous Chloride—A mixture of 1 gramme of Zinc Oxide and 3 c c of Stannous Chloride T S should not assume a dark colour in the course of an hour, *PG*.

Preparation

UNGUENTUM ZINCI ZINC OINTMENT

Add 3 of finely sifted Zinc Oxide gradually to 17 of Benzoated Lard, previously melted at a low temperature, stir until cold.
(1 in 6½)

Official in Austr, 15 in 100, Belg, Dan, Dutch, Fl, Ger, Hung, Ital, Jap, Mex, Norw, Russ, Span, Swed and Swiss, 1 in 10, US, 1 in 5.

Not Official

EMPLASTRUM ZINCI OXIDI (Aseptic) —Zinc Oxide, 20, Resin, 15, Japan Wax, 4, Benzoated Beef Tallow, 25, Anhydrous Wool Fat, 15, Washed Rubber, 8, Glycerin, 12, Methyl Salicylate, 0.6, Thymol, 0.4, all by weight — *Y P B '07*, 429, *C D '07*, n 178, *P J '07*, n 125

GELATINUM ZINCI DURUM (*Unna*) —Dissolve Gelatin 15 and Glycerin 25 in Water 45 Rub down Zinc Oxide 10 with Glycerin 15, mix, and add sufficient Water to produce 100 All by weight

Gelatinum Zinci —Dissolve Gelatin 6 in Distilled Water 18, rub down Zinc Oxide 4 with Glycerin 11, add the Gelatin solution, and mix thoroughly —*B P C Formulary* 1901

GLYCÉRÉ D'OXYDE DE ZINC —Zinc Oxide, 1, Glycerin of Starch, 2 —*F*

LASSAR'S PASTE —Zinc Oxide, 24, Starch, 24, Salicylic Acid, 2, Soft Paraffin, 50 Used in eczema

An unusual case of poisoning by absorption from use of Lassar's paste — *L '04*, 1 432

This has been incorporated in the *B P C* under the title *Pasta Zinci Composita*

PASTA UNNA —Gelatin, 15, Zinc Oxide, 10, Glycerin, 30, Water, 40 Melt, stir carefully, then add Ichthyol (Ammon) 2 p c —*King's*

Pasta Zinci et Ichthamolis —Zinc Oxide, 10, Ammonium Ichthyo-sulphonate, 2, Gelatin, 16, Glycerin, 32, Distilled Water, *q s* to produce 100 — *B P C*

PESSUS ZINCI —Zinc Oxide, 15 grains, Mass (Glyco-gelatin), 20 grains — *Women's*

Pessus Zinci Oxidi —Zinc Oxide, 15 grains, Oil of Theobroma, to 120 grains —*B P C*

PILULA ZINCI ET BELLADONNÆ —Zinc Oxide, 2 grains, Extract of Belladonna (*B P '85*), $\frac{1}{2}$ grain, Extract of Gentian, *q s* —*Charming Cross*

Pilula Zinci Oxidi et Belladonnæ —Zinc Oxide, 2 grains, Alcoholic Extract of Belladonna, $\frac{1}{2}$ grain —*St Thomas's*

This has been incorporated in the *B P C*

PULVIS ZINCI OXIDI COMPOSITUS *Syn* Dusting Powder —Zinc Oxide, 3, Salicylic Acid, in fine powder, 1, Starch, 12 —*Squire*

Pulvis Zinci Oxidi et Acidi Salicylici —Zinc Oxide, 4, Salicylic Acid, in fine powder, 1, Starch, 15 —*B P C*

PULVIS ZINCI OXIDI ET ACIDI BORICI —Zinc Oxide, Boric Acid, in powder, equal parts —*St Thomas's*

This has been incorporated in the *B P C*

PULVIS ZINCI OXIDI ET AMYLI —Zinc Oxide, 1, Starch Powder, 1 — *St Thomas's*

This has been incorporated in the *B P C*

PULVIS ZINCI ET CALOMELANOS —Zinc Oxide, Mercurous Chloride, Tannic Acid, and Starch, of each 1 —*Westminster*

PULVIS ZINCI OLEATIS COMPOSITUS (*Squire*) —Zinc Oleate, in fine powder, 20, Boric Acid, in fine powder, 70, finely powdered French Chalk, 10

UNGUENTUM ZINCI CUM ACIDO SALICYLICO —Salicylic Acid, 20 grains, Zinc Ointment, $\frac{1}{2}$ oz, Soft Paraffin, $\frac{1}{2}$ oz —*Middlesex*

UNGUENTUM ZINCI STEARATIS —Zinc Stearate, 50, White Petrolatum, 50 To the White Petrolatum, melted on a water-bath, add the Zinc Stearate, continue the heat until smooth, then stir while cooling, until it congeals —*U S P*

This has been incorporated in the *B P C*

ZINC OLEAS (Shoemaker's) — Dissolve 180 grains of Zinc Acetate in 40 fl oz of cold Water, add slowly 20 fl oz of a Solution of Sodium Oleate, made by dissolving powdered Castile Soap, 1 oz in 20 fl oz of Water, wash the precipitate with cold Water, collect and dry

It forms a solid cake, easily powdered, and melting at about 79.4°C (175°F)

Solution of Sodium Oleate of the above strength is also used to precipitate Bismuth, Copper, and Lead Oleates

ZINC OXIDE PLASTER MULLS (*Unna*) — Containing $\frac{1}{2}$ grain and 1 grain to the sq in

ZINC AND SALICYLIC PLASTER MULL (*Unna*) — Containing Zinc Oxide $\frac{1}{2}$ grain and Salicylic Acid $\frac{1}{4}$ grain to the sq in

ZINC GELATIN (*Unna*) — Zinc Oxide, 10, Gelatin, 10, Glycerin, 20, Water 20

This has been incorporated in the *B.P.C.*, under the title **Pasta Zinc et Gelatini**, giving the quantities respectively, 15, 15, 35, 35

Not Official

ZINCI PERMANGANAS

Reddish purple, crystalline, hygroscopic masses

Solubility — About 1 in 3 of Water, generally with a slight residue

As an injection in chronic urethritis, 1 grain in 8 fl oz of Water — *B.M.J.* '89, i 1458

Not Official

ZINCI PHOSPHIDUM

Minutely crystalline, friable fragments, or a greyish black powder, containing about 24 p c of Phosphorus, corresponding to the formula Zn_3P_2

Solubility — Insoluble in Water or Alcohol (90 p c) Soluble in acids with evolution of Phosphuretted Hydrogen, which is not spontaneously inflammable

Medicinal Properties — Strongly recommended as a substitute for Phosphorus

In hay fever — *P.* lv 205, *P.J.* '95, ii 205

Dose — $\frac{1}{16}$ to $\frac{1}{4}$ grain = 0.0032 to 0.0162 gramme, given in pill with Milk Sugar and Glucose

Foreign Pharmacopœias — Official in Fr (Phosphure de Zinc), Mex and Span

ZINCI SULPHAS.

ZINC SULPHATE

$\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$, eq 285.41

FR, SULFATE DE ZINC OFFICINAI, GER, ZINKSULFAT, ITAL, SOLFATO DI ZINCO, SPAN, SULTATO ZINCICO

Colourless, transparent, somewhat efflorescent, rhombic crystals, white acicular crystals or a granular crystalline powder. It should be kept in well closed bottles

Solubility — 10 in 7 of Water Insoluble in Alcohol (90 p c)

Medicinal Properties — Astringent, given with doubtful result in chorea, also in infantile diarrhoea, in large doses a prompt

emetic As an astringent **injection** in leucorrhœa and in the less acute stages of gonorrhœa, as a **collyrium** in conjunctivitis

Dose.—1 to 3 grains = 0.06 to 0.2 gramme as a tonic, as an emetic, 10 to 30 grains = 0.65 to 2 grammes

Ph. Ger. maximum single dose, 1.0 gramme

Prescribing Notes.—*Tincture or Wine of Opium causes no precipitate with*

Incompatibles of Zinc salts are—Alkalis and their Carbonates, Lime Water, astringent vegetable Infusions or Decoctions and Milk

Antidotes.—In case of poisoning with the salts of Zinc, Sodium Carbonate or Potassium Carbonate in large quantities dissolved in warm Water, Milk and Eggs freely, Tannic Acid or strong Tea, Laudanum, Linseed Meal Poultices to abdomen. If there is much pain in the abdomen, an enema of Guel, or Starch and Water may be given.—*Murrell*

Official Preparations.—Used in the preparation of Unguentum Zinci Oleatis, Zinci Carbonas, and Zinci Valerianas

Not Official.—Bagnarium Zinci Sulphatis, Injectio Zinci Sulphatis, Lotio Rubra, Lotio Zinci Sulphatis, Collyrium de Zinci, and Cadmi Sulphas

Foreign Pharmacopœias.—Official in Austri, Belg, Dan, Dutch, Fl, Ger, Hung, Ital, Jap, Mex, Norw, Port, Russ, Span, Swed, Swiss and U S

Tests.—Zinc Sulphate melts when heated rapidly. At a temperature of 50° C (122° F) it loses 5 molecules of its Water of crystallisation, equivalent to 31.3 p.c., at 100° C (212° F) the same molecule, equivalent to an additional loss of 6.3 p.c. or a total loss of 37.6 p.c. takes place, at a temperature of about 240° C (464° F) it parts with the remaining molecule of Water of crystallisation, equivalent to an additional loss of 6.3 p.c., or a total loss of 43.9 p.c. It dissolves readily in Water, forming a clear solution which is acid in reaction towards blue Litmus paper, and which yields the tests distinctive of Zinc given under that heading. The aqueous solution affords on the addition of Barium Chloride Solution a white precipitate insoluble in Hydrochloric Acid. Neither the *B.P.* nor the *P.G.* states a requisite percentage of pure crystallised Zinc Sulphate, nor is a method of determination indicated. The *U.S.P.* requires that in an unefflorescent condition it should contain not less than 99.5 p.c. of pure crystallised Zinc Sulphate, but gives no method of determination. The impurities mentioned under Zinc should be absent from the Sulphate. It should respond to the tests given in small type below under the headings of Litmus, Sodium Hydroxide, Silver Nitrate, Sulphuric Acid and Ferrous Sulphate, indicating the absence of free acid, Ammonium salts, Chlorides and Nitrates. A standard has been suggested (*C.D.* '08, 1797) of 0.05 p.c. for Chloride, calculated as Zinc Chloride ($ZnCl_2$)

Litmus.—If 2 grammes of Zinc Sulphate be shaken with 10 c.c. of Alcohol (90 p.c.) and, after 10 minutes, filtered, the filtrate, diluted with 10 c.c. of Water, should not affect blue Litmus paper, *P.G.* If a gramme in small fragments be agitated with 10 c.c. of Alcohol (91.9 p.c.) for some time and filtered, the filtrate should not redén in moistened Litmus paper.

Sodium Hydroxide.—Zinc Sulphate should not evolve Ammonia on the addition of 0.1 g. of Sodium Hydroxide, *P.G.*

Hydrogen Sulphide—The aqueous solution of Zinc Sulphate (1-20), after being acidulated with Hydrochloric Acid, should not respond to the time-limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted, *U S P*

Silver Nitrate—The aqueous solution (1-20) should not be rendered turbid by T S of Silver Nitrate, *P G*, not more than slightly turbid, *U S P*

Sulphuric Acid and Ferrous Sulphate—2 cc of an aqueous solution (1-10) of the salt, with 2 cc of Sulphuric Acid added and 1 cc of Ferrous Sulphate T S poured on as a layer, should not give a coloured zone, even on standing for some time, *P G*

Preparation

UNGUENTUM ZINCI OLEATIS ZINC OLFATE OINTMENT

Precipitate a solution containing 2 of Zinc Sulphate in 4 of Distilled Water with a solution of Hard Soap 4 in Distilled Water 40, wash the precipitated Oleate with hot Distilled Water until free from Sulphate, dry and mix with an equal weight of the Soft Paraffin, melted, stir until cold

The Zinc Oleate is now made by precipitation instead of dissolving Zinc Oxide in Oleic Acid

Not Official

BUGINARIUM ZINCI SULPHATIS—Zinc Sulphate, $\frac{1}{10}$ grain, Oil of Theobroma, 40 grains — *Westminster*

INJECTIO SULPHATUM—Zinc Sulphate, Copper Sulphate, Ferrous Sulphate and Alum, of each 1 grain, Water to 1 fl oz — *Lock Hospital*

This has been incorporated in the *B P C*

INJECTIO ZINCI SULPHATIS—Zinc Sulphate, 3 grains, Water 1 fl oz

For gonorrhoea and leucorrhoea

This has been incorporated in the *B P C*, as follows —

Zinc Sulphate, 0.75, Distilled Water, *q s* to produce 100

LOTIO RUBRA—Zinc Sulphate, 2 grains, Compound Tincture of Lavender, 10 minims, Water, to 1 fl oz A stimulant to indolent ulcers

This has been incorporated in the *B P C*, as follows —

Zinc Sulphate, 0.50, Compound Tincture of Lavender, 2, Distilled Water, *q s* to produce 100

LOTIO ZINCI SULPHATIS—Zinc Sulphate, 1 grain, Distilled Water, 1 fl oz Used in ophthalmia — *London Ophthalmic*

COLLYRE AU SULFATE DE ZINC—Zinc Sulphate, 0.15, Rose Water, 100 — *Fr*

Antiseptin is stated to be a mixture of Zinc Sulphate and Iodide, Thymol and Boric Acid

Zinc Sulphide (Zinc Sulphite) is a white crystalline powder, sparingly soluble in Water It has been used as a relatively non toxic antiseptic for impregnating gauze and dressings

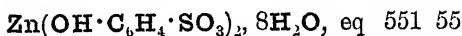
CADMI SULPHAS—Colourless crystals, readily soluble in Water, in soluble in Alcohol Has been used as an astringent in the place of Zinc Sulphate

Official in Fr, Mex and Port

ZINCI SULPHOCARBOLAS.

ZINC SULPHOCARBOLATE.

ZINC PHENOL-PARA-SULPHONATE



Colourless or almost colourless efflorescent rhombic crystals, sometimes possessing a faint Phenol odour and

The *B P* gives the formula for Zinc 1 molecule of Water of crystallisation. It is officially stated that it may be obtained by heating a mixture of Phenol and Sulphuric Acid, and saturating the product with Zinc Oxide. It has been pointed out in subsequent editions of the *Companion* that, prepared in this way, it will contain a quantity of Sulphate. The salt contains 8 molecules of Water of crystallisation. The *U S P* describes the salt under the heading Zinc Phenolsulphonas or Zinc Phenolsulphonate.

It should be kept in well-closed glass bottles of a dark amber tint and exposed as little as possible to the air, as the salt effloresces and has a tendency to become pink on exposure to air.

It is extraordinary that the official monograph should describe the salt as 'efflorescent' when the official formula shows only 1 molecule of Water of crystallisation, as a matter of fact, the commercial salt contains 8 molecules of Water of crystallisation, and corresponds to the above formula.

Solubility—1 in 2 of Water, 3 in 1 of boiling Water, 1 in $2\frac{1}{2}$ of Alcohol (90 p c).

Greenish and Smith (*P J* '02, 1 552) give the solubility of the salt as 1 in 2.7 of Water and state that on consideration it is thought that Zinc Phenol-para-sulphonate would be fairly stated to the fact that it is a Para salt. The experiments made to determine whether the salt yielded a constant weight when dried at a given temperature not being satisfactory, the solubility was therefore determined gravimetrically by precipitation with Sodium Carbonate as usual and the result calculated for a salt of the official formula, ignoring, moreover, the fact that an amount of Zinc Oxide corresponding to even the 'assumed' official formula had not been obtained. The process would have worked admirably had the official formula been correct, but Squire and Cames (*C D* '02, 11 945) have shown that the reason for their inability to obtain a constant weight was the incorrectness of their assumption of the pharmacopoeial formula, the salt containing 8 molecules of Water of crystallisation and not 1. Greenish and Smith's figure (1 in 2.7) for Zinc Sulphocarbolic as it should properly be understood, is incorrect. Their subsequent statement (*P J* '03 1 917) that the correct formula for a salt of this composition is $\text{Zn}(\text{OHC}_6\text{H}_4\text{SO}_3)_2, 11\text{H}_2\text{O}$. If calculated figures are to be admitted as 'authoritative,' the percentage of Magnesium in, say, Magnesium Sulphate might with equal reason be determined by the usual method, and the solubility calculated for a salt containing 1 molecule of Water of crystallisation instead of 7, provided the official volume was sufficiently condescending to adopt an incorrect formula.

for the salt By a curious oversight a misprint occurs in their second note on the solubility of Zinc Sulphocarbolate and Ammonium Phosphate In giving their final conclusion that if the composition of the official salt is altered from 1 molecule of Water of crystallisation to 8, then the solubility of the salt must be proportionately increased, they have given the formula for the Mono-hydrated Zinc Sulphocarbolate as $\text{Zn}(\text{OHC}_6\text{H}_4\text{SO}_3)_2 \cdot \text{HO}_2$, instead of $\text{Zn}(\text{OHC}_6\text{H}_4\text{SO}_3)_2 \cdot \text{H}_2\text{O}$ The criticisms referring to the solubility figure for Ammonium Phosphate appear under the heading of Ammonium Phosphate

Medicinal Properties —Astringent and antiseptic

For a spray to the throat, 5 grains to the oz of Water, for a nasal douche, 2 grains to the oz of Water, for vaginal injection, 60 grains in a pint of Water, for leucorrhœa or gonorrhœa

Foreign Pharmacopœias —Official in Austri and Jap (Zincum Sulphocarbolium), Dutch (Sulphophenylas Zincicus), Russ and Swiss (Zincum Sulfophenicum), US (Zinc Phenolsulphonas)

Tests —Zinc Sulphocarbolate when heated to 100°C (212°F) loses 6 molecules of Water of crystallisation, equivalent to a loss of 19.45 p c At 125°C (257°F) it loses the remaining molecules of Water of crystallisation, equivalent to an additional loss of 6.48 p c, or total loss of 25.93 p c When more strongly heated it chars, evolves an odour of Phenol and leaves a residue amounting to about 14.6 p c of the original weight It dissolves readily in Water, forming a solution which is acid in reaction towards blue Litmus paper, and which then diluted yields with Ferric Chloride T S a violet colour and which answers the tests distinctive of Zinc given under that heading The *BP* neither states a requisite percentage nor a method of determination The *USP* requires that the uneffloresced crystals should contain not less than 99.5 p c of pure Zinc Paraphenol sulphonate, but gives no method of determination It may be determined gravimetrically by precipitating a weighed quantity of the salt, dissolved in a definite volume of Water, by the addition of Ammonium Carbonate Solution, filtering off the precipitated Zinc Carbonate, washing, drying, igniting and weighing as Zinc Oxide 80.8 parts of Zinc Oxide correspond to 408.5 parts of anhydrous Zinc Sulphocarbolate or 426.39 parts of Zinc Sulphocarbolate of the *BP* formula with 1 molecule of Water of crystallisation, or to 551.55 parts of crystallised Zinc Sulphocarbolate of the correct formula with 8 molecules of Water of crystallisation The impurities mentioned under Zinc should be absent from this salt It should conform to the tests given below under the headings of Barium Chloride, Silver Nitrate, and modified Gutzeit's test, indicating the absence of Sulphates, Chlorides, and Arsenic

Time-limit Test —The aqueous solution of the salt (1–20) to which 1 c c of diluted Hydrochloric Acid has been added, should not respond to the time limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted, *USP*

Barium Chloride —The aqueous solution of the salt (1–20) should not become turbid upon the addition of T S of Barium Chloride, *USP*

Silver Nitrate.—A similar solution should not become turbid upon the addition of 1% of Silver Nitrate, *USP*

Gutzeit's Test.—5 cc of an aqueous solution of the salt (1-10) should not respond to the modified Gutzeit's test for Arsenic, *USP*.

ZINCI VALERIANAS.

ZINC VALERIANATE

ZINC-ISO-VALERIANATE



FR, VALERIANATE DE ZINC, GER, ZINKVALERIANAT, ITAL, VALERIANATO DI ZINCO, SPAN, VALFRIANATO ZINCICO

White lustrous pearly scales, having a strong odour of Valerianic Acid and a sweetish metallic taste

It should be kept in well-stoppered bottles, as when exposed to the air it slowly loses Valerianic Acid. The *BP* gives the formula for the anhydrous salt, the salt really contains 2 molecules of Water of crystallisation, the *USP* gives the formula with 2 molecules of Water of crystallisation. The official method of preparation is by saturating Iso-valerianic Acid with Zinc Carbonate, or by the reaction of Zinc Sulphate and Sodium Iso-valerianate

Solubility—1 in 120 of Water, 1 in 60 of Alcohol (90 p c), 1 in 500 of Ether

Medicinal Properties.—Antispasmodic and nervine tonic, used in various neuralgic and hysterical affections, and sometimes in chorea

In hay fever —*BMJ* '96, 1967

Dose—1 to 3 grains = 0.06 to 0.2 gramme

Incompatibles—All acids, soluble Carbonates, most metallic salts and vegetable astringents

Not Official.—Pilula Valerianæ Composita, and Pilula Zinci Valerianatis

Foreign Pharmacopœias—Official in Dutch, Fr, Hung, Ital, Jap, Mex, Port, Russ, Span, Swed and U S

Tests.—Zinc Valerianate when heated melts, and at a higher temperature is decomposed, giving off vapours, and leaving a residue of Zinc Oxide which when dissolved in Diluted Hydrochloric Acid should answer the tests distinctive of Zinc given under that heading. It dissolves sparingly in Water, forming a solution which possesses an acid reaction towards blue Litmus paper. The *USP* states that it should dissolve without residue in Ammonium Carbonate. 0.5 of a gramme of the salt when dissolved in a mixture of 0.5 cc of Hydrochloric Acid and 9 cc of Water gives a liquid from which the Iso-valerianic Acid separates and floats as an oily layer. The *BP* does not state a requisite percentage of pure Zinc Valerianate, but gives a method of determination, stating that it should yield not less than 26 nor more than 30 p c of Zinc Oxide. The theoretical percentage of Zinc Oxide is 26.8, in the salt containing 2 molecules

of Water of crystallisation, in the salt of the present official formula it is 30.4 p.c. A number of commercial samples yielded from 21 to 64 p.c. of Oxide and suggested a minimum standard of 26 p.c. All the samples examined showed Butyric Acid by the Copper test. The commercial 'precip' generally contains a quantity of Oxide, but pure samples can occasionally be obtained.

The *U.S.P.* states that the salt should contain not less than 99 p.c. of pure Zinc Valerate ($2\text{H}_2\text{O}$), but gives no method of determination. The salt should not contain the impurities mentioned under Zinc. When testing for Butyric Acid the *B.P.* tests the distillate with Copper Acetate Solution, the *U.S.P.* tests the concentrated aqueous solution of the salt with a concentrated Copper Acetate Solution by the test described in small type below under the heading of Copper Acetate. The salt should answer the tests given in small type below under the headings of Silver Nitrate, Barium Chloride, Ferric Chloride and modified Gutzzeit's test.

Time-limit Test—If 0.5 gramme of the salt be dissolved in a mixture of 0.5 c.c. of Hydrochloric Acid and 9 c.c. of Water, the Valeric (Iso Valeric) Acid will be liberated and float as an oily layer on the surface of the liquid. After filtering through a wetted filter, the clear solution should not respond to the time-limit test for Arsenic, Cadmium, Lead and Copper, in applying this test the addition of Ammonia Water should be omitted, *U.S.P.*

Barium Chloride—If 0.5 gramme be dissolved in a mixture of 0.5 c.c. of Nitric Acid and 4.5 c.c. of Distilled Water and the mixture filtered through a small wetted filter, the filtrate should show but a faint cloudiness upon the addition of T.S. of Barium Chloride, *U.S.P.*

Silver Nitrate—A similar filtrate should show but a faint cloudiness upon the addition of T.S. of Silver Nitrate, *U.S.P.*

Ferric Chloride—If 0.5 gramme be triturated with 3 c.c. of Water, and 0.2 c.c. of Ferric Chloride T.S. added, the filtrate should not show a red colour, *U.S.P.*

Copper Acetate—This T.S. should not immediately affect the transparency of the distillate obtained on heating the salt with Diluted Sulphuric Acid. Only drops are formed after the lapse of a little time, and these gradually pass into a bluish white crystalline deposit, *B.P.* A mixture of a concentrated solution of Copper Acetate in Water and a concentrated aqueous solution of the salt should remain perfectly clear, *U.S.P.*

Gutzzeit's Test—If 0.5 gramme of the salt be heated with a mixture of 9.5 c.c. of Distilled Water and 0.5 c.c. of Hydrochloric Acid and filtered the filtrate should not respond to the modified Gutzzeit's test for Arsenic, *U.S.P.*

Not Official.

PILULA VALERIANÆ COMPOSITA—Zinc Valerianate, Iron Valerianate and Quinine Valerianate, of each 1 grain—*Samaritan*.

This has been incorporated in the *B.P.C.* under the title **Pilulæ Ferri Valerianatis Compositæ**, *syn* **Pilulæ Trium Valerianatum**.

PILULA ZINCI VALERIANATIS—Zinc Valerianate, 1 grain, Compound Pill of Asafetida, 2 grains—*Throat and St. Thomas's*.

This has been incorporated in the *B.P.C.*

ZINGIBER.

GINGER

FR, GINGEMBRE, GER, INGWER, ITAL, ZENZERO, SPAN, RIZOMA DE JENGIBRE

The scraped and dried Rhizome of *Zingiber officinale*, Roscoe
From plants cultivated in the West Indies, India, and other countries

Medicinal Properties.—Aromatic stimulant and carminative
It is given in atonic dyspepsia, flatulence, and as a corrective adjunct to purgative medicines

Official Preparations—Syrupus Zingiberis, and Tinctura Zingiberis, used in the preparation of Infusum Sennæ, Pilula Scillæ Composita, Pulvis Cinnamon Compositus, Pulvis Jalapæ Compositus, Pulvis Opii Compositus, Pulvis Scammonii Compositus Contained in Mistura Sennæ Aloes et Ferri, and Pilula Camphoræ Composita The Tincture is used in the Acidum Sulphuricum Aromaticum, Liquor Sennæ Concentratus Composita, and contained in Infusum Cinchonæ Acidum

Not Official—Tinctura Zingiberis Fortior, and Oleoresina Zingiberis

Foreign Pharmacopœias—Official in all, Fr, Gingembre, Ital, Zenzero, Port, Gengibre, Mex and Span, Jengibre

Descriptive Notes.—The rhizome of Ginger comes into commerce in several forms, which are either coated, unscraped, or entirely scraped, and in some varieties washed with lime. The last are termed bleached.

The rhizome differs in being either starchy and brittle with a fibrous fracture, or hard and resinous and rather tough, also in the degree of pungency.

The bleached Jamaica Ginger is considered to be the best for flavour, and the Cochin next, that of Fiji, which is rare in commerce, has a characteristic lemon flavour. The West African Ginger, although inferior in appearance and in smaller pieces, is often superior in pungency. The pieces, which are known technically as pieces of hands, are in the finer Jamaica and Cochin varieties blanched laterally, about 3 in (75 mm) long, the branches being compressed, more or less oval, and contracted below, and at the rounded end exhibiting a depression corresponding to the base of the leafy stem. The scraped surface is of a pale buff colour and fibrous appearance, the taste hot and pungent and the flavour characteristic.

In some specimens of Ginger the appearance is honeycombed, owing to the root having been scalded before drying, but usually it is mealy towards the apices, even when resinous below. Its odour is due to $\frac{1}{4}$ p.c. of a volatile Oil, but its pungency to an oily body named Gingerol.

An inferior variety known as Batoon Ginger is sometimes imported from the West Indies, it consists of the younger shoots, which are generally kept for propagating the plant. Japanese Ginger is occasionally imported, it has a greyish fracture, occurs in smaller pieces and is apparently derived from a different species.

Powdered Ginger is characterised by the cells containing resin, by

the pyriform compressed starch grains 12 to 40 μ long, appearing linear when seen laterally, having the hilum at the smaller end, by the thin-walled polygonal parenchymatous cells, and the sometimes septate bast fibres with irregularly nodose extremities

Tests—Ginger yields when genuine and unbleached from 3 to 4 p c of ash, and 5 p c should not be exceeded. It yields not less than 1.5 p c of soluble ash. The extractive matter soluble in cold Water is usually about 10 p c, and should not be less than 8.5 p c, the extractive matter soluble in Alcohol (90 p c) usually amounts to about 5 p c, and should not be materially below this figure

Preparations

SYRUPUS ZINGIBERIS SYRUP OF GINGER

1 of Ginger (in the form of strong Tincture 1 in 2), Syrup 7 s to yield 40 (1 of Ginger in 40)

Dose— $\frac{1}{2}$ to 1 fl dm = 1.8 to 3.6 c c

Official in Jap, 1 of Tincture in 10, Swed, 1 (rhizome) in 20, by weight, U S, 3 (Fluid Extract) in 100

TINCTURA ZINGIBERIS TINCTURE OF GINGER

1 of Ginger in No 40 powder, percolated with Alcohol (90 p c) to yield 10 (1 in 10)

Dose— $\frac{1}{2}$ to 1 fl dm = 1.8 to 3.6 c c

Official in Belg, Ger, Hung, Ital, Jap, Mex, Port, Swiss and U S, 1 in 5, all by weight except U S

Tests—Tincture of Ginger has a sp gr of 0.835 to 0.840, it contains about 0.5 p c w/v of total solids and about 88 p c w/v of Absolute Alcohol

Not Official

TINCTURA ZINGIBERIS FORTIOR *Syn* Essence of Ginger (D P '85)

—Ginger percolated with Alcohol (90 p c) to form 1 in 2

Dose—5 to 20 minims = 0.3 to 1.2 c c

Squire's **Essence of Ginger** has always been twice the above strength

By repercolation a Fluid Extract 1 in 1, or even 2 in 1, can be readily prepared

U S has Fluidextractum Zingiberis 1 in 1 with Alcohol (95 p c)

OLEORESINA ZINGIBERIS (U S) *Syn* Gingerine

Ginger, in No 60 powder, exhausted by percolation with Acetone and evaporation

Should be kept in a well stoppered bottle

CINCHONÆ RUBRÆ CORTEX.

QUINQUINA ROUGE (*F*)

DETERMINATION OF TOTAL ALKALOIDS AND QUININE

Process of the *French Codex* (1908)

The amount of total alkaloids and crystallised basic quinine of Water of crystallisation, which the *Codex* is required to yield, are briefly referred to on p. 382 under the comparison of the respective quantities of the more important This section of the work was too far advanced to enable the to be introduced under the heading of Cinchona, and it is included here in view of the interest attaching to the subject

Tests—The time available for a trial of the undermentio necessarily been short, but so far as it is possible to judge, except involved in manipulating such large quantities of solvents of so volatile and inflammable a nature, the process works well. If the solvents are not subsequently recovered by distillation, the process is not economical. Details of the process are as follows. A sample of the bark, 50 grammes, is powdered and the powder passed through a No. 45 sieve. The amount of moisture is determined on a weighed quantity of 0.5 of a gramme, drying the powder at 100° C (212° F). The bark under examination lost at this temperature moisture equivalent to 8.6 p.c., so that 100 parts of the original powdered bark may be considered as equivalent to 91.4 parts of the dried and powdered bark. A weighed quantity of 30 grammes of the dried and powdered bark is introduced into a wide mouth bottle, and a previously prepared mixture of 85 c.c. of Ammonium Hydroxide (10 p.c. w/v) and a sufficient quantity of Alcohol (95 p.c.) to produce a total capacity of 125 c.c. is poured upon it. The mixture is allowed to stand for 1 hour, and then time to time and 720 c.c. of Ether then added. The stopper of the bottle is securely tied in by the aid of string passed round the neck of the bottle, the contents are briskly shaken and allowed to remain at rest for 6 hours with intervals of frequent shaking, the liquid is filtered through a plated filter paper contained in a covered funnel, and a measured quantity of 750 c.c. of the solution (= 25 times of the dried and powdered bark) is removed. The whole of the Ether is distilled, evaporation being conducted several times in a flask of 500 c.c. capacity and away from the naked flames, the flask being simply plunged into warm Water. After the Ether is completely distilled the evaporation is continued until a portion of the Alcohol is also removed, the liquid is transferred to a flask of 125 c.c. capacity and the distillation continued until nothing further passes over. The volatile Alkaloids are removed by immersing the flask up to the neck in a water-bath, the residue which remains is dissolved in Water, solution being effected by gently warming the mixture, and then allowing the solution to cool when the acid solution is added. An unplated filter and collected in a glass separator of 250 c.c. capacity. A measured quantity of 125 c.c. of Chloroform is introduced, followed by a sufficient quantity of dilute Ammonia Solution to liberate the alkaloids and to produce a distinct ammoniacal odour, the addition of the Ammonia produces a considerable reddish-brown deposit, colouring matter, etc., which holds a large amount of the Chloroform in suspension necessitating filtration through a glass filter under pressure, and the washing of this precipitate to free it from retained alkaloids. The chloroformic solution of the alkaloids is separated, referred to a flask and the treatment or the ammoniacal solution in the separator twice repeated, using in each instance 125 c.c. of Chloroform. The mixed chloroformic solutions are separated in each instance, washed with 10 c.c. of Water, allowed to separate and the aqueous washings rejected. The chloroformic solution is distilled in such a flask as at 200 c.c. of liquid remain, the chloroformic liquid is transferred, after cooling, to a

flask graduated at 250 c c. The flask in which the distillation has been conducted is washed with a few small quantities of Chloroform, and the washings transferred to the graduated flask and diluted with sufficient Chloroform to bring the volume to 250 c c, they are then thoroughly mixed. A measured quantity of 50 c c (= 5 grammes of the dried and powdered Cinchona bark) is removed and evaporated to dryness in a conical flask of 90 c c capacity, the residue is dried at 100° C (212° F) and weighed, the weight of the residue multiplied by 20 yields the amount of alkaloids contained in 100 grammes of powdered Cinchona bark, and this weight should not be less than 5 grammes. In the experiment under consideration the amount of total alkaloid amounted to 8.8 p c and was fairly highly coloured. When treated with Normal Volumetric Hydrochloric Acid Solution, using Hematoxylin Solution as an indicator of neutrality and calculating the result with a factor for anhydrous Quinine it indicated 5.78 p c of alkaloids.

Determination of Quinine.—The Chloroform is removed by distillation from the remaining 200 c c of the above Chloroform solution of the total alkaloids, the syrupy residue is treated with 50 c c of Ether which is dropped on a little at a time, the Cinchonine and the greater part of the Cinchonidine will be precipitated in a crystalline condition. When the precipitate has settled down, the ethereal solution is decanted into a flask, the crystals washed by decantation with 75 c c of Ether, used in 3 portions, the ethereal solutions of Quinine in the flask are mixed, almost the whole of the Ether distilled and the liquor so concentrated is transferred to an evaporating dish and allowed to evaporate spontaneously, the Ether washings are also transferred to the dish previous to evaporation. The almost colourless sticky residue is dissolved in 20 c c of Sulphuric Acid Solution (2 p c w/w) by warming on a water bath until solution is completed, dilute Ammonia Solution is added little by little to the warm limpid solution until the precipitate at first formed ceases to redissolve. The liquid will now be slightly turbid and alkaline, and 5 p c w/w Sulphuric Acid Solution is added carefully drop by drop until the liquid, now rendered limpid, yields a very faintly acid reaction towards blue Litmus paper, the volume should be about 15 c c which should be cooled and allowed to crystallise during 12 hours in a cool place, the crystals collected on a porcelain dish of 20 mm diameter pierced with a hole which should be covered with a circle of flannel with a diameter a little bit larger than the hole, the complete apparatus being placed in a very small filter, which has been previously moistened with Distilled Water. The crystalline Sulphate is transferred to the funnel washed with 6 c c of Distilled Water, used in 3 separate portions and which have previously served to wash out the evaporating basin in which the crystallisation was effected. The funnel is inverted over a piece of white filter paper, the basic Quinine Sulphate in the shape of a compact cake is detached and dried in the air, when the drying has proceeded sufficiently far the circle of flannel is separated, which will be found to be cleanly effected without retention of the product, the cake of Sulphate is placed in a watch glass, at the same time transferring any particles which may have adhered to the flannel, and complete desiccation is effected at a temperature of 100° C (212° F) until of constant weight. The Sulphate should be weighed between 2 watch glasses held together by means of a metal clip, the whole having been previously weighed. The weight of Sulphate obtained should not be less than 0.251 gramme, which corresponds to 1.257 grammes of basic Quinine Sulphate (dried at 100° C (212° F)), or to 1.092 grammes of anhydrous Quinine in 100 grammes of the dried and powdered bark. This is equivalent to a yield of not less than 1.5 p c of basic Quinine Sulphate ($C_{20}H_{21}N_2O_2$)₂ · H₂SO₄ · 8H₂O, to 1.257 p c of basic Quinine Sulphate dried at 100° C (212° F), or to 1.092 p c of anhydrous Quinine from the dried and powdered bark. The crystallised basic Sulphate of Quinine obtained in the experiment in question was of good crystalline appearance, and was practically free from colour. The percentages calculated on the dried and powdered bark corresponded to 4.08 p c of basic Quinine Sulphate ($C_{20}H_{21}N_2O_2$)₂ · H₂SO₄ · 8H₂O, eq. to 3.425 p c of basic Quinine Sulphate dried at 100° C (212° F), or to 2.97 p c of anhydrous Quinine.

Not Official

THERAPEUTIC AGENTS OF BACTERIAL ORIGIN.

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THERAPEUTIC SERA

Syn—ANTITOXINS, OR ANTI-SERA

These are obtained by treating an animal with subcutaneous or intravenous injections of increasing doses of (a) bacterial toxins, (b) bacterial cultures, living or killed, (c) a combination of a and b, then bleeding the animal, allowing the blood to coagulate, drawing off the serum and bottling this in the form of ampoules after drying *in vacuo*, all these operations being carried out under aseptic precautions. To the fluid serum a small quantity of an antiseptic is usually added, and each bottle or vial generally contains a single dose only. The dried serum should be in the form of thin scales or fine powder, otherwise it is difficult to dissolve, for use each gramme (corresponding to about 10 c.c. of fluid serum) of the solid is dissolved in 5 to 10 c.c. of cool Distilled Water (not above 40° C = 104° F) previously sterilised by boiling.

Two classes of anti-sera may be distinguished, one prepared by method a, with bacterial toxins, to which the term 'antitoxin' is alone strictly applicable (e.g., diphtheria and tetanus antitoxins), the other prepared by method b, and termed anti-microbial, or simply anti-sera (e.g., anti-streptococcus, anti-plague, and anti-pneumonic sera).

The last named are much less potent than the antitoxins, and attempts have been made to reinforce their action by the simultaneous injection of fresh normal serum, but without much success.

It is customary in some instances to employ several strains of the organism in the preparation of the serum, such sera are termed 'polyvalent'. The subject of serum treatment is fully dealt with in Hewlett's 'Serum Therapy'.

The therapeutic sera in most instances retain their activity for several weeks at least if kept in a cool, dark place—preferably an ice-safe. Diphtheria and tetanus antitoxins probably for nearly a year, the anti-microbial sera for a much shorter period. They should not be administered with any other substance, must not be heated, and a bottle of the fluid having once been opened, any fluid not used at the time should be discarded. The dried products are preferable in hot climates.

The dose of antitoxin corresponds usually to 5 to 20 c.c. of the fluid serum, according to the gravity of the case, which is estimated by ascertaining the amount of serum required to neutralise a given amount of toxin or culture. The dose depends on the gravity of the disease and not on the age of the patient. Some estimation on general principles should be employed in addition to the above.

The therapeutic sera are administered by subcutaneous injection, in the abdomen or between the scapulae, the skin having been previously disinfected with an antiseptic lotion and the syringe by boiling for five minutes. Or, if an immediate effect be desired, by intravenous injection into a large vein, the serum being warmed by standing the bottle in warm Water (not above 40° C = 95° to 104° F) and strained through a piece of fine muslin sterilised by boiling, if there be any deposit. Care must be taken that no air is injected. Intra-muscular injections are more quickly absorbed than subcutaneous ones. Early treatment is of the utmost importance.

Some clinicians assert that anti-sera exert their action when

administered by the mouth or rectum. Hewlett, however, was unable to detect any absorption of tetanus antitoxin when given in this way to rabbits, and Sternberg similarly no absorption of diphtheria antitoxin — *Wien klin Wochr* 1908, p 709

The therapeutic sera are specific, e.g., diphtheria antitoxin is of use only in diphtheria, carefully administered they are harmless, but cutaneous eruptions or joint pains may follow, for the treatment of which Calcium Chloride is of service

A second injection of serum at an interval of 10 to 40 days after the first one may be followed by immediate and serious symptoms ('supersensitisation,' see Goodall, *Jour of Hyg* vii 1907, p 607) But the continuous use of a serum for some days does not produce this effect

The anti sera may be used as prophylactics (dose 10 to 20 c.c. subcutaneously), but the immunity produced does not last longer than three weeks

DIPHTHERIA ANTITOXIN — Anti diphtheritic serum is official in the *Ph. Ger.* Both a liquid and a solid antitoxin are described, the former as a yellowish, transparent fluid, having the odour of the preservative agent, and with at most a slight sediment, the latter is a yellowish white powder, or yellow transparent lamellæ, which, by the addition of 10 parts of Water, dissolves to a liquid corresponding in colour and general appearance to the liquid diphtheria antitoxin

The sizes of the liquid diphtheria antitoxin mostly used are No 0, 200 immunisation units, No 1, 500 to 600 immunisation units, No 2, 1000 immunisation units, No 3, 1500 immunisation units

The solid diphtheria antitoxin is required to contain at least 5000 immunisation units per gramme

The *Ph. Ger.* also stipulates that antitoxin, with marked permanent turbidity or thick deposit, as well as serum of a prohibited test number, is not permitted to be sold in pharmacy

It should be protected from the light and stored in a cool place

The therapeutic value is reckoned in Ehrlich units, 1 unit being that amount of serum which will completely neutralise about 100 lethal doses of toxin in a medium sized guinea pig

The method of standardisation is a very exact one, devised by Ehrlich, but is too complicated to explain here. The different makes are of different strengths, i.e., contain a variable number of units in a given volume. The dosage is always referred to in units

The dosage varies with the severity of the attack, and with the lapse of time after the onset before treatment is commenced. In a mild case, coming under observation on the first day, a single dose of 4000 units may suffice, but is best repeated on the next day. In severe cases the 4000 units should be repeated every 4 hours for 3 or 4 doses, and repeated the next day if necessary. In bad cases, coming under observation late, 8000 to 30,000 units have been recommended, followed by smaller doses, every 3 or 4 hours. In such cases Cairns considers that valuable time is saved by giving the primary dose intravenously. If there be a reasonable suspicion that the case is diphtheritic no time should be lost in giving antitoxin. The guide to the administration of subsequent doses is the general condition and the appearance of the membrane, this when the patient is fully under the influence of antitoxin appears to melt away

The prophylactic dose should not be less than 500 units

Several preparations by different makers can be obtained

Diphtheria antitoxin, *re* the intravenous injection, even in cases other than laryngeal, it seems difficult to say (*L. '04*, ii 1776) of an individual patient that a better result was obtained by this method than might have followed subcutaneous injection

TETANUS ANTITOXIN — If the case be seen immediately upon the development of the premonitory symptoms (stiffness, etc., of the facial muscles), 25 to 30 c.c. of the serum may be injected subcutaneously, followed by an injection of 10 c.c. every 8 hours as long as the symptoms last. If any time has elapsed since the development of the premonitory symptoms, 10 c.c. should be administered intravenously and 20 c.c. subcutaneously, followed by 10 c.c.

subcutaneously every 8 hours as before. But if the case has lasted any length of time, and especially if symptoms have already occurred, no time should be lost in giving the antitoxin by cerebral or spinous inoculation (*see infra*).

Dried and pulverised tetanus antitoxin has been recommended as a dressing for wounds soiled with earth, etc.

For prophylactic use 3 doses of 10 c c should be injected at intervals of a fortnight.

In veterinary practice, 20 to 40 c c may be injected every 12 to 24 hours, but unless the animal be a valuable one the cost of the treatment is prohibitive.

Since Tetanus Toxin becomes fixed in the cells of the central nervous system, and antitoxin is but slowly absorbed from the subcutaneous tissues, it is desirable, in order to obtain a maximum and rapid response, to inject the antitoxin so that it may at once come in contact with the nerve tissues. This may be done by (a) intra-cerebral, into the cerebral hemispheres, (b) into the lateral ventricles, (c) intra-spinous, by primary dose of 5 c c of tetanus antitoxin of 1 gramme of the solid dissolved in 5 c c of Water may thus be administered, subsequent doses being given subcutaneously. As tetanus toxin is absorbed along the nerve trunks, antitoxin may be injected in addition into nerves if the site of the wound permit.

A method given by Roux and Borrel — *T G '08*, 773

A method given by Semple — *B M J '99*, 1 10

10 c c doses, given intradurally in a case of tetanus, were followed by recovery — *B M J '04*, 11 1696

ANTI-VENENE — An antitoxin prepared by the injection of snake venom. A separate serum is required for every venom, so that this antitoxin must have a limited use. That prepared by Calmette, of Lille, is mainly antidotal for the venom of the cobra. At least 30 to 40 c c should be injected at the earliest possible moment if any interval has elapsed since the bite, 10 c c should be given intravenously in addition.

Antivenomous sera have been shown (*L '04*, 11 1277) to be markedly if not absolutely specific, even between the venoms of species of the same genus. The only sera at present in practical use are Calmette's and one prepared at the Pasteur Institute of India with pure cobra venom. Both are specific for cobra venom. The neutralising power is low, and 300 to 400 c c may be necessary even when given intravenously, and 10 or 20 times this amount if given subcutaneously.

ANTI-STREPTOCOCCIC SERUM — The dose is 10 to 20 c c every 12 or 24 hours. Some Continental authorities regard this amount as much too small, and administer 50 to 150 c c for a dose.

N B — Streptococcic serum rapidly diminishes in strength with age, and should not be kept.

Is of especial value in erysipelas. Some cases of septicæmia react well to it, in others apparently similar it has little effect. Cases of septicæmia may be due to a variety of organisms, but it is only in pure streptococcic infections that the serum can be expected to have any effect. Even in streptococcic infections it is not always efficacious, there seem to be many varieties of streptococci, and a serum prepared with one variety may have little or no antidotal action towards another variety. The serum should be a 'polyvalent' one, *z c*, prepared with several varieties or strains.

The use of a serum prepared from a horse which has been immunised against a variety of strains has been recommended (*L '04*, 11 1829), the commencing dose being at least 20 c c, if necessary, at least every 24 hours. Unless to persist in the use of any particular serum unless its beneficial action is almost immediately apparent.

The serum is innocuous if carefully prepared and injected with due precaution. (2) It must be administered early in the disease, and in large doses — 20 c c — twice in 24 hours in severe cases. (3) If administered early and in large doses definite improvement is observed in a considerable proportion of cases — *B M J '05* 1 584

The most rational method of dosage (*L* '04, n 1832) would seem to be that of a large injection on the first occasion, followed by smaller doses as case may require. More uniform results are obtained with polyvalent sera.

In a simple septicæmia or sapræmia good results can be obtained with the anti streptococcic serum — *L* '04, n 1213

Fenwick recommends rectal injections of polyvalent anti-streptococcic serum in gonorrhœa, gonorrhœal pyæmia, rheumatism and hæmorrhagic purpura — *B M J* '06, i 979

A prophylactic injection recommended by Cheyne previous to operations about mouth and throat

ANTI-ANTHRAX SERUM — Prepared by immunising asses with killed and living cultures of *B anthracis*. Sclavo's is that generally employed, dose 50 c c

A successful case, *B M J* '05, n 118

ANTI-PNEUMOCOCCIC SERUM — The dose is 20 to 30 c c subcutaneously twice daily until the crisis. If the case be seen *early*, this serum may be very useful in the case of debilitated, aged, or alcoholic patients. Pane's serum seems to be the most potent.

Washbourn, *B M J* '97, i 510, n 1849, and Eyre, *ib* '99, n 1247, Wilson, *Jour Amer Med Assoc* 1900 (Sep), 595, Tyler, *ib* 1901 (June), 1540

ANTI-PLAGUE SERUM — Yersin's serum is that generally employed. Calmette recommends 20 c c intravenously to be given immediately, followed by two subcutaneous doses of at least 40 c c each during the first 24 hours, and subsequently 10 to 40 c c daily, according to the condition of the patient. Choksy and also Cairns recommend still larger doses (60, 80, 100, 200 c c). The prophylactic dose is 10 c c subcutaneously.

Calmette, *Ann de l'Inst Pasteur*, xiii '99, 865, Cairns, *L* '08, i 1287

ANTI-TYPHOID SERUM — No satisfactory serum seems to have been prepared as yet. The sera on the market are anti-microbic, dose 10 to 20 c c. Chantemesse has prepared a serum, the use of which he claims gives good results.

Chantemesse, *La Presse Méd* '02, No 103, 122, *Trans XIV Internat Congress Hygiene*, Macfadyen, *B M J* '03, i 681

Anti-typhoid serum is stated (*B M J* '04, n 1269) to exercise a specific action on the diseases of the organism. If the nervous system is deeply poisoned, the benefit is much smaller, and failures occur. In the treatment of a large number of cases during several years, Chantemesse shows a mortality of 4 p c., and claims (*B M J* '04, n 1449) to have produced a serum with which remarkable success is stated to have been obtained. The serum is given in small doses (4 to 5 m), and Wright believes that it contains a *toxin* and is in reality a *vaccine* (Chantemesse agrees with this view). In the case of typhoid fever (*B M J* '04, n 1449), the serum of a horse, after repeated inoculations with the virus, though possessed of anti-bacterial properties, is found to be practically devoid of any antitoxic value.

ANTI-TYPHOID EXTRACT OF JEZ — Prepared from the tissues of immunised rabbits. Dose, 2 dm, by the mouth every 1 to 2 hours until temperature becomes remittant.

See *B M J E* '01, i 51, '02, i 27

ANTI-TUBERCLE SERUM — Paquin and Maragham have each prepared an anti-tubercle serum.

Dr Marmorek has obtained a new tuberculous serum by growing the young bacilli in a medium consisting of leucotoxin calf serum and Glycerin liver bouillon. In pulmonary tuberculosis he claims, by the use of this antitoxin, to have produced amelioration, and even definite cures. In pleurisy there was a rapid diminution of the effusion — *B M J* '03, n 1434, '06, i 340, *L* '08, n 1470 1612, 1746

Not very favourably reported on — *B M J* '01, n 1621, *L* '03, n 1695

The dose for rather chronic cases is ordinarily 5 c.c., whilst in acute cases, such as meningitis, as much as from 20 to 30 c.c. in divided doses may be given every day for 4 or 5 days, the dose being then gradually diminished. The serum when given in carefully graduated doses, with proper precautions and in suitable cases, does no harm. Experience tends to show that the serum does produce a specific antitoxic effect.—*L '04*, 1 859, 979, *B M J '04*, 1 749, 857.

The treatment has so far proved rather disappointing (*L '04*, 11 1827), Marmorek's serum proving no more successful in active and progressive cases than other sera. The method of administration recommended in the *Edinburgh Medical Journal*, 1905, 213, is 3 c.c. injected on the first day, 4 c.c. on the second, 5 c.c. on the third day, no injections for the next three days, 5 c.c. on the seventh, 6 c.c. on the eighth, 7 c.c. on the ninth, and 8 c.c. on the tenth day. This completes the first series, and an interval of eight to ten days is allowed. Then 8 c.c., and in another series of eight injections the amount is raised to 20 c.c. Another interval, and then a further series of injections similar to the latter.

An interesting and instructive lecture on Antitoxins was delivered by Professor R. Tannei Hewlett at an evening meeting of the Pharmaceutical Society, and is duly reported in the *P J '04*, 11 888, and *C D '04*, 11 975.

An interesting résumé of the tuberculin and anti-tuberculous sera is given (*L '05*, 1 923), the conclusion being that, judging from evidence, the use of tuberculin materially improves the results of treatment, and it would seem quite justifiable to supplement sanatorium methods with this specific one.

The method now recommended by Marmorek is the injection of 5 c.c. every other day for 3 weeks, followed by a clear 3 weeks' interval, after which the injections are repeated as before. The site of injection is preferably the abdominal walls or thighs, and should be varied as much as possible.

(1) The beneficial effects of the serum are most marked in the 'surgical' forms of the disease.

(2) No objectionable features whatsoever follow the injections.

(3) The pyrexia is diminished, though the fall in temperature is sometimes preceded by an initial rise.

(4) Pain in the 'surgical' forms is almost invariably alleviated.

(5) The expectoration of sputum in pulmonary tuberculosis is rapidly diminished.

(6) The patient's health rapidly improves and the patient gains in weight.

—*L '05*, 1 777.

Cases on record (*L '05*, 11 603) which would seem to show that it undoubtedly exerts a neutralising effect on the tuberculosis toxin. In all cases in which the serum may be tried it is advisable not to push it unduly, and sufficient intervals should be allowed between the injections.

A valuable paper (*L '05*, 1 928), by H. Batty Shaw, on the treatment of tuberculosis of the lungs by means of tuberculin and other bacterial derivatives. A history of the different tuberculin is given and their respective doses. Dr. Marmorek's serum is again referred to, and a review of the work of various experimenters is recorded. The general conclusion seems to be that, judging from the evidence, it would seem that the use of the tuberculin materially improves the results of treatment. Tuberculin treatment is of little value, but may be justifiable to supplement the ordinary treatment by the latter methods.

Many anti-tuberculous serums have been produced and others have attracted some attention. The results so far achieved have been inconclusive and, for the most part, disappointing.—*B M J '05*, 1 1393.

SERUM FOR HAY FEVER—Dunbar, by injecting horses with the toxin extracted from the pollen of various *Gramineae*, has obtained an anti-serum which is stated entirely to allay the troublesome symptoms of hay fever. The fluid serum is applied frequently to the eyes, a solid powder to the nose. The remedy is sold under the name of 'Pollantin'.

A lengthy paper appears in the *American Journal of Pharmacy*, 1905, 1 1, giving a résumé of this suggested remedy for hay fever. It is prepared in powder and in liquid form. The method of using is as follows:

(1) Pour about a third of the contents of the serum-phial into the accompanying empty glass-phial, provided with a dropping pipette. The latter is dropped in

sent out in a small wooden case, and should be carried in the pocket as nearly as possible in the upright position

(2) The method to employ in using liquid Pollantin is as follows (a) For the eye Bring, by means of the pipette, one drop to the outer angle of the eye, and, drawing down the lower lid with the finger, allow the drop to come into contact with the mucous membrane. A pleasantly cool sensation felt in the eye shows that the instillation has been properly carried out

(b) For the nose With the head bent somewhat backwards, insert the point of the pipette about $\frac{1}{2}$ inch into each nostril, and express one or two drops of Pollantin into each. Care must be taken to keep the pipette squeezed so long as it is within the nose, otherwise the Pollantin will be drawn back into the pipette again. After Pollantin has been introduced into one nostril the other must be kept closed while the serum is snuffed up from the one treated, tapping the while on the outside of that nostril with the finger

(3) The pipette, together with the india rubber head, should be thoroughly cleansed at least once daily, and kept for one minute in boiling water

The powder is obtained (I J P '05, 335) by completely drying the serum *in vacuo* at 45° C (113° F) and mixing it with sterilised Milk Sugar. It forms a yellowish and almost colourless powder. This should be snuffed into the nostrils or blown in with an insufflator, and can be dusted upon the conjunctiva with a camel's hair brush

The method of using the powder —(1) A portion of the pulverised Pollantin as large as a lentil is dropped into the little scoop attached to the stopper of the bottle. The scoop is then held under one of the nostrils, the other nostril being compressed and occluded by the finger. The powder is then snuffed into the open nostril, the snuffing being repeated several times, during which the ala of the nostril is lightly tapped with the finger to distribute the powder over as much of the mucous membrane as possible. (2) If the powder is also to be used for the eyes, the accompanying camel's hair brush is lightly dipped into it, the brush being then gently applied to the inner surface of the attached lower lid, or a small quantity of the powder may be shaken upon it from the brush. With each new bottle of the powder a new brush should also be brought into use

Distressing symptoms following the use of 'Pollantin', giddiness, tinnitus, vertigo and vomiting —L '05, n 130

Semon, *B M J* '03, i 713, '04, i 1168

CANCER SERUM —Various sera have been prepared for malignant growths. A recent one is that of Schmidt, but reports of its use are not encouraging —L '03, n 1374, *B M J* '04, i 299

Doyen has reported the discovery of a micrococcus (*M. neoformans*) in cancerous tumours, and with the organism has prepared an anti serum. The results of the inquiry at the Pasteur Institute and elsewhere are to establish that the *M. neoformans* does sometimes exist in cancerous tumours. Dudgeon finds that it is almost identical with *M. pyogenes albus* (*Jour Hyg* vii '07, 13). As regards the serum, the tumours have diminished in size, but as to whether the serum can prevent recurrence, requires some years to answer —*B M J* '04, n 1712, *L* '04, n 1799

It is stated (*B M J* '05, n 211) that during the five months in which all the cases which M. Doyen has wished to show have been examined, no single case of amelioration has been seen. Records of a further series of cases treated with the serum also appear in *L* '06, i 955, 1186, but in no case was any benefit seen

NORMAL SERUM FOR ALIMENTATION —Normal horse serum, heated to 60° C (140° F) for half an hour, may be used (a) to replace for a short time, or (b) to supplement, gastric or rectal feeding in cases of vomiting, obstruction, etc. For (a) children, 30 to 50 cc, adults 100 to 150 cc, for (b) 20 to 40 cc should be given subcutaneously daily

Many other anti sera have been prepared, but are of doubtful value and are not on the market (e.g., cholera, dysentery, hydrophobia, leprosy, scarlatina, syphilis, whooping-cough, etc.).

Serum injections, e.g., horse serum and diphtheria antitoxin, sometimes produce good effects in many diseases, e.g., arthritis, gonorrhoea, arthritis, asthma, broncho pneumonia

TUBERCULIN PREPARATIONS.

A—KOCH'S ORIGINAL TUBERCULIN—Prepared by boiling, concentrating, and filtering three months-old Glycerin broth cultures of the tubercle bacillus

An amber-coloured, syrupy fluid, with a characteristic odour. Gives the reactions for Glycerin and for albumoses

The maximum initial dose should not exceed 0.001 c.c. and is administered by subcutaneous injection. The injection is followed in tubercular subjects by a rise of temperature of 2° to 5° F., and constitutional disturbance more or less severe. The dose must not be repeated until the reaction produced by the preceding one has completely passed off. The same dose is administered until it is followed by only a slight reaction, a larger amount may then be given, increasing by 0.001 c.c. until 0.005 is reached, then by 0.002 c.c. and so on.—Watson Cheyne, *Med. Chirurgical Trans.* 1891.

Goetsch considers that it is undesirable to obtain a reaction, and therefore commences with very small doses, 0.00001 to 0.0001 gramme, if even the former produces reaction treatment is commenced with the *new* tuberculin 0.001 increasing to 0.1 milligramme. When this is reached treatment is continued with the old tuberculin commencing with 0.0001 to 0.001 gramme.

For diagnostic purposes the initial dose should not exceed 0.005 c.c., which if it produces a reaction, may be followed by 0.01 c.c. and 0.02 c.c. Tuberculin for cutaneous reaction has been applied to the skin after scarification (Von Pirquet's cutaneous reaction). Precipitated with Alcohol and the precipitate dissolved in the solution used in Calmette's test.

(For the diagnosis of tuberculosis 0.001 c.c. to 0.02 c.c.)

For the commencement of treatment a 1 p.c. solution is used, later on a 10 p.c. solution. The dilutions should be made in a solution of Carbolic Acid, and only so long as the reaction is present. Now almost discarded for treatment.

Tuberculinum Kochi is included in the *Ph. Ger.* It is described as a clear, light brown fluid, possessing a pleasant aromatic odour. It is readily miscible with Water. It contains in addition to the active constituents about 40 parts of Glycerin in 100 parts, as well as the constituents of the bouillon, but no antiseptics.

It is put up in flasks bearing an official leaden seal, and only the undiluted preparation is allowed to be held in stock. The dilutions recommended by the physician are directed in all cases to be freshly prepared, and sterilised. Distilled Water or, still better, a 0.5 p.c. Carbolic Solution to be used in the preparation thereof. It must be kept in a cool place and protected from the light.

The tuberculins possess distinct value as a specific means of treatment. An interesting and encouraging account of the results which have followed the use of the original tuberculin of Koch was given (*B. M. J.* '05, i, 1393) by Professor McCullagh at an address at the annual meeting of the Dermatological Society of Great Britain and Ireland. The treatment extends over a period of six or more months. Beginning with an initial dose of from $\frac{1}{4}$ to $\frac{1}{2}$ c.c. of a 1 in 1000 solution, the amount is gradually increased, according to the constitutional reaction obtained, up to as much as 1 c.c. of pure tuberculin. The injections are repeated every third or fourth day. Old tuberculin is still a valuable remedy, capable of producing satisfactory and even brilliant results.

Old tuberculin is not trustworthy in intra-ocular tuberculosis and it has done much harm. Useful as a means of diagnosis in intra-ocular tuberculosis, but useless in treatment.—*B. M. J.* '05, ii, 432.

Not necessary to employ tuberculin in gradually increased doses for diagnostic purposes. Procedure consisted in administering a small dose of $\frac{1}{10}$ milligramme of Koch's old tuberculin in adults and $\frac{1}{20}$ milligramme in children, repeating the injection if necessary after an interval of 3 or 4 days.—*L.* '05, ii, 1203.

B—KOCH'S NEW TUBERCULIN—There are three varieties, termed respectively A, O and R. Tuberculin R or T R, is the only one of therapeutic value.

It is prepared by titrating and emulsifying virulent tubercle bacilli with Distilled Water and centrifugalising. The fluid contains 2 milligrammes of solid matter per c.c. (not 10 milligrammes as formerly stated, *B M J* '08, 1 468).

The fluid is administered by subcutaneous injection after diluting with sterile 20 p.c. Glycerin Solution. The preliminary doses should correspond to not more than $\frac{1}{100}$ of a milligramme of solid matter, i.e., 0.5 c.c. of a dilution of 1:500. The doses now recommended are usually $\frac{1}{10000}$ to $\frac{1}{3000}$ milligramme administered every 10 to 14 days and controlled by opsonic determinations.

The impossibility of limiting the reactive energy of tuberculin to the skin prevents, in many cases, the utilisation of specific properties in the treatment of lupus vulgaris.—*B M J* '05, 1 689.

The results gained by injections of Tuberculin T.R. are at least as good as those by any other method. It is not assumed that in Tuberculin T.R. a perfect remedy for tuberculosis of the urinary system exists, but for vesical tuberculosis it seems the best remedy at our disposal.—*L* '05, 11 1769, *B M J* '05, 11 1587.

Tuberculin R given in doses of about $\frac{1}{10000}$ milligramme at intervals determined by estimations of the opsonic power of the blood is a most valuable weapon in the fight against pulmonary tuberculosis.—*B M J* '06, 11 18.

Latham has adduced clinical and bacteriological evidence that Tuberculin R and bacterial vaccines produce their therapeutic effects when administered by the mouth.—*Proc. Roy. Soc. Med.* 1 '08, Med. Sec. 195.

REFERENCES.—*L* '97, 11 568, 600, 704, 1488 '98, 11 194, *B M J* '97, 11 207, '98, 1 357, '98, 11 77, *B M J E* '97, 11 19, 27, 31, 55, 103, '98, 1 47, 55, *T G* '97, 850, '98, 400, *P* lix 399. Oxy Tuberculin.—*L* '98, 1 179, *B M J E* '98, 11 27.

OPSONINS

Great interest has been aroused by the discovery of the significance attached to the opsonic power of the blood and the value of estimations of the opsonic index in the diagnosis and treatment of tuberculosis and other infections. The term 'opsonin' was invented by Wright and is derived from a classical word which means 'to cater or prepare victuals for,' and it apparently prepares the bacteria for ingestion by the phagocytes. Opsonins are described (*L* '05, 11 1917, *B M J* '05, 11 842) as substances contained in the serum or plasma of blood which possess the power of so modifying various kinds of bacteria as to render them an easier prey to the attacks of leucocytes.

This opsonic power is found in the blood of both healthy and diseased persons, but differs in degree, an essential difference being that whereas the degree is held to be approximately the same in normal healthy persons, wide variations are found amongst those who are diseased. The process of determining the opsonic index of the blood in tuberculosis is briefly outlined (*B M J* '05, 11 172) as follows.—Equal quantities of the patient's serum, an emulsion of tubercle bacilli (or other organisms), and leucocytes (washed in a solution of $\frac{1}{2}$ p.c. Sodium Citrate in normal salt), are taken in a capillary pipette and incubated together for 15 minutes, after which films are made of the mixture and stained in a modified way for tubercle bacilli, etc., the number of tubercle bacilli (or other organisms) ingested by 50 polynuclear white corpuscles is counted, and the figure thus obtained is compared with a standard similarly obtained, but using the serum of a healthy person. The former figure divided by the latter gives the 'opsonic index'. A series of experiments on the 'opsonin' treatment of tuberculous patients were undertaken at the London Hospital Medical College, and the results are recorded in the *L* '05, 11 1603.

An address on the opsonic theory and its practical application to medicine and surgery is reported in the *B M J* '06, 11 16, and gives a very lucid and up to date review of the present knowledge of this subject. (See also *Practitioner*, May 1908, Wright and others.)

VACCINES.

Vaccines are used either for prevention or prophylaxis, or for the treatment of chronic or sub-acute infections. For the latter, cultures of the organism corresponding to, and preferably isolated from, the infection, are sterilised by

heat and standardised as to the number of organisms they contain in a given volume from which the dose is calculated. For tuberculous infection Tuberculin R is used (see above).

GLYCERINATED VACCINE LYMPH is prepared by mixing calf lymph with 50 p.c. of Glycerin and storing for three months, this destroys all extraneous organisms. A preparation in which the extraneous organisms are killed with Chloroform is also prepared (Green's method) (Prophylactic)

LEPROLIN.—A toxin of the cultivation of the *Bacillus lepro*. It is stated (B M J '05, 1 699) to have given good results in the treatment of leprosy.

By incubating leprosy tissue in salt solution Deycke obtains a growth of an acid-fast streptothrix. A fatty substance extracted from this (termed Nastin) used for the treatment of leprosy.—B M J '08, 1 802

STAPHYLOCOCCIC VACCINE is prepared from cultures of the *S. pyogenes*, *aureus* and *albus* (dose 100–1000 millions). Valuable for locococic infections, e.g., acne, furunculosis, sycosis, etc.—B M J

CHOLERA VACCINE is prepared from virulent cultures of the cholera spirillum. Dose, 1 c.c. (Wright and Bruce, B M J '93, 1 227) (Prophylactic)

TYPHOID VACCINE is prepared from virulent cultures of the *Bacillus typhosus* (Wright and Semple, B M J '97, 256, Wright and Leishman, ib '00, 1 122) (Prophylactic)

PLAGUE VACCINE is prepared from cultures of the *Bacillus pestis* (B M J '97, 1 1057 and 1461) (Prophylactic)

Streptococcic (dose, 20–60 millions), Pneumococcic (dose, 10–50 millions), Gonococcic (dose 100–500 millions), Malta fever, *Bacillus coli* (dose 5–10 millions) and other vaccines have been used with success in the controlling chronic or sub-acute infections. Vaccine treatment should be controlled by opsonic determinations.

ANTI-RABIC INOCULATION.—The Pasteur system of inoculation as practised for bites of rabid animals. Emulsions of spinal cord of rabbits dried for periods varying from 14 to 3 days and injected subcutaneously. The treatment must be carried out at an Institute (e.g., Pasteur Institute, Rue Dutot, Paris, Pasteur Institute, Lille) (Useless for treatment of disease when declared)

Some of the anti-sera (diphtheria, tetanus, plague, and streptococcus) may be employed as prophylactics, but their protective power is transient (three weeks), whereas the vaccines protect for at least many months.

COLEY'S FLUID

A fluid prepared by cultivating the streptococcus of erysipelas and the *Bacillus prodigiosus* in broth, and heating to 55°C (136°F) for one hour.

It has been used in the treatment of malignant growths, especially sarcomata. The dose to commence with is $\frac{1}{2}$ to 1 minim, administered by injection in the neighbourhood of the tumour. The dose is gradually increased, the guide being the amount of reaction produced.

MALLEIN

Prepared by boiling and concentrating both cultures of the glanders bacillus and the mallein, for the diagnosis of glanders in animals. The requisite dose is injected into the neck. In a glandered animal a large swelling forms at the seat of the injection, the local lesion becomes enlarged, and the temperature rises at the site of the injection. It is of no therapeutic value.

DE BACKER'S FLUID

Pure cultures of yeast stored under pressure in syphon-like vessels provided with hollow needles by means of which the dose is injected.

Has been used in the treatment of tuberculosis and cancer.—B M J '97, 1 802

CHEMICALS REAGENTS, ETC.,

USED IN QUALITATIVE TESTING

In the undermentioned list of substances and solutions used in chemical analysis, the word 'parts,' where it relates to preparations of the German Pharmacopœia, is to be understood to mean 'parts by weight'

ACIDUM ACETICUM ACETIC ACID

BP } The acids official in the respective Pharmacopœias
USP }
PG —The *PG* Acidum Aceticum Dilutum more closely resembles Acetic Acid *BP* and *USP* See Acidum Aceticum, p 10

ACIDUM ACETICUM GLACIALE

BP } The Glacial Acetic Acid official in the respective Pharmacopœias
USP } See Acidum Aceticum Glaciale, p 12
PG —Acidum Aceticum *PG* is practically Glacial

ALBUMEN

BP —A thin glairy liquid obtained from the egg of *Gallus Bankiva* var *domesticus*, from which the shell and yellow yolk have been separated

USP } Not included
PG }

Cogulated White of Egg is employed by the *BP*, *USP* and *PG* as a test for the activity of Pepsin

ALBUMEN SOLUTION

BP —Albumen, 1 cc, Water, 4 cc The solution is recommended to be freshly prepared, and either the above quantity of Water may be added or a sufficiency to adjust the solution to meet the requirements of individual tests The solution may be strained through moist cotton wool or tow

USP —The liquid portion of a fresh hen's egg, freed from the yolk, mixed with 100 cc of Water and filtered It is recommended that the solution be prepared fresh for use when required

PG —Not included

ALCOHOL, ABSOLUTE

BP }
PG } The Absolute Alcohol official in the respective Pharmacopœias
USP }

ALCOHOL, DILUTED

BP —The *BP* employs 90 pc and 70 pc Alcohols, the *USP* uses four strengths, viz, the Alcohol of the Pharmacopœia (94.9 pc), 90 pc, 80 pc, and 70 pc, and the *PG* uses the official 'Spiritus' containing from 90 to 91.2 pc of Ethylic Alcohol All by volume

ALUM.

B P—A colourless, transparent crystalline salt represented by the chemical formula $\text{Al}_2(\text{SO}_4)_3$, K_2SO_4 , $24\text{H}_2\text{O}$, or $\text{Al}_2(\text{SO}_4)_3$, $(\text{NH}_4)_2\text{SO}_4$, $24\text{H}_2\text{O}$

P G } Not included
U S P }

AMMONIA SOLUTION

B P—The official 'Liquor Ammoniae'

P G—The official 'Liquor Ammonii Caustici'

U S P—The official 'Aqua Ammoniae'

They each contain 10 p c by weight of Ammonia For strong solution of Ammonia the *B P* directs the use of the official 'Liquor Ammoniae Fortis'

AMMONIUM ACETATE SOLUTION

B P—The official 'Liquor Ammonii Acetatis'

P G } Not included
U S P }

AMMONIUM CARBONATE SOLUTION

B P—A filtered solution of 5 grammes of Ammonium Carbonate, 7.5 cc of Ammonia Solution, and Distilled Water *q s* to yield 100 cc

P G—Ammonium Carbonate 1 part, dissolved in a mixture of Water 3 parts and Solution of Ammonia 1 part

U S P—Ammonium Carbonas, *U S P* 20 grammes dissolved in a mixture of Ammonia Water 20 cc and Water 70 cc, adding *q s* of Water to measure 100 cc

AMMONIUM CHLORIDE SOLUTION

B P } A filtered solution of 10 grammes of Ammonium Chloride in Dis-
U S P } tilled Water *q s* to measure 100 cc

P G—Dissolve 1 part of Ammonium Chloride in 9 parts of Water

AMMONIUM CHLORIDE, NESSLER'S SOLUTION OF

B P—A solution of 0.315 gramme of Ammonium Chloride in recently boiled and Ammonia free Distilled Water *q s* to measure 100 cc

AMMONIUM CITRATE SOLUTION

B P—The official 'Liquor Ammonii Citratis'

P G } Not included
U S P }

AMMONIUM HYDROSULPHIDE See AMMONIUM SULPHIDE

AMMONIUM MOLYBDATE

B P—A white or almost white crystalline solid, represented by the chemical formula $(\text{NH}_4)_2\text{MoO}_4$

U S P } Not included
P G }

AMMONIUM MOLYBDATE SOLUTION

B P—Ammonium Molybdate, 10 grammes, Water, *q s* to produce 100 cc, filter

U S P—Ammonium Molybdate 15 grammes, Ammonia Solution, if necessary, *q s* to effect 100 cc This solution is gradually poured into a volume of Nitric Acid [sp gr 1.403 at 25°C (77°F)] to a gentle heat for about 2 hours, and decanted from any yellow deposit which may form An alternative method is to mix gradually and with repeated shaking a solution obtained by dissolving Molybdic Acid (H_2MoO_4) 10 grammes, Ammonia Solution, 42 cc, with a solution obtained by mixing 63 cc of Nitric Acid, of the above sp gr with an equal volume of Water Heat gently for two hours Any yellow sediment

separating out after the solution has been made some days may be separated by decanting the liquid in the same manner as above. It is recommended that the reagent be kept in the dark, and the clear solution decanted from any sediment which may separate out from time to time.

P G—Not included

AMMONIUM OXALATE

B P—The colourless crystalline salt represented by the chemical formula $(\text{NH}_4)_2\text{C}_2\text{O}_4 \cdot \text{H}_2\text{O}$

P G
U S P } Not included

AMMONIUM OXALATE SOLUTION

B P—A filtered solution of 2.5 grammes of Ammonium Oxalate in Distilled Water *q s* to measure 100 c.c.

P G—A solution of 1 part of Ammonium Oxalate in 24 parts of Water

U S P—The solution of 4 grammes of pure crystallised Ammonium Oxalate $[(\text{NH}_4)_2\text{C}_2\text{O}_4 \cdot \text{H}_2\text{O}]$ in Water *q s* to measure 100 c.c. An alternative method is to dissolve 4 grammes of pure Oxalic Acid in 100 c.c. of Water, add 15 c.c. of Ammonia Water, boil to expel excess of Ammonia, and dilute with Water to 113 c.c. The *U S P* requires that the residue obtained on evaporating a portion of the solution should, when ignited, be completely volatilised. The absence of Chlorides and Sulphates should be proved by the precipitate produced by Silver Nitrate T.S. or by Barium Chloride T.S. being completely soluble on the addition of Nitric Acid.

AMMONIUM SULPHATE

B P
P G } Not included

U S P—The salt represented by the chemical formula $(\text{NH}_4)_2\text{SO}_4$ and may be prepared by neutralising a mixture of equal volumes of pure Sulphuric Acid and Water with Ammonia Water, then evaporating the solution and crystallising. The alkalinity of the liquid should be maintained during the evaporation, if necessary, by the addition of more Ammonia, ascertained by testing from time to time with Litmus paper. Three grammes of the salt should leave no appreciable residue upon ignition. An aqueous solution of the salt (1-10) should not respond to the time limit test for heavy metals, nor should it become turbid with Nitric Acid and Silver Nitrate T.S. A similar solution of the salt should not be coloured red by 2 drops of Hydrochloric Acid and 1 drop of Ferric Chloride T.S. (absence of Sulphocyanate).

AMMONIUM SULPHIDE SOLUTION

B P—The *B P* uses a solution of Ammonium Hydrosulphide prepared by passing washed Hydrogen Sulphide through 60 c.c. of Solution of Ammonia until saturated, then adding a further 40 c.c. of Solution of Ammonia. It recommends that the solution be prepared when required.

P G—Not included

U S P—The solution is prepared by saturating 3 parts of Ammonia Water with Hydrogen Sulphide, and then converting the greater portion of the Ammonium Hydrogen Sulphide formed into Ammonium Sulphide by the addition of 2 parts of Solution of Ammonia. It should be kept in small dark amber-coloured bottles in a cool, dark place, and when a notable deposit of Sulphur has made its appearance the solution should be rejected. It should be a perfectly clear and colourless solution, leaving no residue on evaporation, nor should any turbidity be produced in it by Magnesium Sulphate T.S. or by Calcium Chloride T.S.

If Ammonium Polysulphide T.S. be required it may be prepared by dissolving a small quantity of pure Sulphur in Ammonium Sulphide T.S.

AMMONIUM THIOCYANATE

B P—The crystalline salt represented by the chemical formula NH_4SCN

P G
U S P } Not included.

AMMONIUM THIOCYANATE SOLUTION

B P—A filtered solution of 2.5 grammes of Ammonium Thiocyanate in Distilled Water *q s* to measure 100 c.c.

P G. }
U S P } Not included

AMYL ALCOHOL

B P—A clear, colourless liquid, sp. gr. about 0.812, which is obtained by fractionating Fusel Oil, after the impurities soluble in a saturated Sodium Chloride Solution have been removed by shaking with that menstruum. The fraction distilling between 125° and 142° C (257° and 289° F) may be reserved for use. It contains principally Iso-amyl Alcohol. The boiling point of Iso-amyl Alcohol at *p* may here be noted, is between 128° to 132° C (262.4° to 269.6° F).

U S P—A colourless oily liquid, boiling at 131° C (267.8° F). Soluble 1 in 40 of Water at 25° C (77° F), miscible with Alcohol *γ*, Ether (sp. gr. 0.716 at 25° C (77° F)), Chloroform, Carbon Disulphide, Petroleum Benzene, Benzene fluid and volatile oils.

P G—A colourless completely volatile liquid (sp. gr. 0.814). Boiling point 129° to 131° C (264.2° to 267.8° F).

AURIC CHLORIDE SOLUTION

B P—A solution containing approximately 4.28 p.c. Aurous Chloride or 3 p.c. w/v of Auric Trichloride prepared by dissolving 1.2 grammes of commercial Gold in a mixture of 1.5 c.c. of Nitric Acid and 1.5 c.c. of Hydrochloric Acid, adding a further 1 c.c. of Hydrochloric Acid, and evaporating to dryness at a temperature of 100° C (212° F) free from acid vapours, and dissolving the residual Chloride in 50 c.c. of Water.

P G—Not included

U S P—Convert commercial Gold Chloride [consisting chiefly of Chlorauric Acid ($\text{HAuCl}_4 \cdot 4\text{H}_2\text{O}$)] into neutral Auric Chloride by fusing it at a temperature not exceeding 150° C (302° F), moistening the residue with enough hot Water to produce a syrupy liquid. The clear liquid poured off from the precipitate and mixed with 20 volumes of Water gives the test-solution. An alternative method is to dissolve 1 gramme of dry Auric Chloride in 30 c.c. of Water.

BARIUM CARBONATE

B P. }
P G. } Not included

U S P—The purified salt represented by the chemical formula BaCO_3 , prepared by precipitating a solution of 12 parts of purified crystallised Barium Chloride in boiling Water 30 parts, with Ammonium Carbonate, 5 parts, followed by Ammonia Water, 5 parts. The precipitate is then thoroughly washed and dried.

BARIUM CHLORIDE

B P—The official salt is in colourless crystals and contains 2 molecules of Water of crystallisation, and is represented by the formula $\text{BaCl}_2 \cdot 2\text{H}_2\text{O}$, eq. 242.54. It is officially required to contain not more than traces of Iron, as indicated by its aqueous solution yielding no precipitate with Ammonium Hydrosulphide Solution, and to be free from alkali and alkali earthy metals as indicated by its failure to leave a residue when the filtrate, after the complete removal of Barium by diluted Sulphuric Acid, is evaporated to dryness in a platinum dish. It will be noted that the *U S P*, which gives a similar test, evaporates to dryness, and heats to redness. Unless the amount of diluted Sulphuric Acid corresponds exactly to that necessary to precipitate the whole of the Barium, a slight residue of the concentrated acid will remain. The *U S P* directions are therefore somewhat more explicit. The *B P* permits the use of either Barium Nitrate, $\text{Ba}(\text{NO}_3)_2$, eq. 259.56, or Barium Acetate, $\text{Ba}(\text{C}_2\text{H}_3\text{O}_2)_2$, eq. 253.56, provided they manifest a similar degree of freedom from these impurities.

U S P—A salt of similar composition to that of the *B P*, but the aqueous solution is required to be perfectly neutral, not to respond to the time-limit test for heavy metals, namely, Antimony, Arsenic, Cadmium, Copper, Iron, Lead and Zinc, to be free from traces of Strontium, as indicated by the colour imparted to a non luminous flame by diluted Alcohol which has been allowed to remain in contact with the salt for some hours, the colour which it is required should be imparted being that of a pure yellowish green free from red, and it is required to be free from other fixed bases as determined by completely removing the Barium by diluted Sulphuric Acid, evaporating the filtrate to dryness and heating on platinum foil

P G—The salt is official in the text of the *P G* See Barium Chloride Not in the list of reagents

BARIUM CHLORIDE SOLUTION

B P—A 10 p c w/v clear filtered solution of the official salt

U S P—A 10 p c w/v solution of the official salt

P G—A 5 p c w/w solution of Barium Nitrate in place of a solution of the Chloride

BARIUM HYDROXIDE

B P—A crystalline salt represented by the chemical formula $\text{Ba}(\text{OH})_2 \cdot 8\text{H}_2\text{O}$, eq. 818 20, prepared by the purification by recrystallisation of the precipitate obtained by the interaction of concentrated Barium Chloride and Sodium Hydroxide Solutions. It is required to be free from heavy metals, e.g., Arsenic, Copper, Iron, Lead and Zinc, as ascertained by its aqueous solution yielding no precipitate with Ammonium Hydrosulphide Solution, and to be free from metals of the alkalis and alkali earths as ascertained by the evaporation to dryness of the filtrate after the complete removal of the Barium as Barium Sulphate by means of a dilute Sulphuric Acid, only a very slight residue should remain

P G } Not included
U S P }

BARIUM HYDROXIDE SOLUTION

B P—A filtered solution of 5 grammes of Barium Hydroxide in recently boiled Distilled Water *q s* to measure 100 c c

P G—A solution of crystalline Barium Hydroxide, 1 part in 19 parts of Water

U S P—A saturated aqueous solution of the salt $\text{Ba}(\text{OH})_2 + 8\text{H}_2\text{O}$, to be prepared when required for use

BARIUM NITRATE SOLUTION

B P—See Barium Chloride

P G—A 5 p c w/w solution of Barium Nitrate

U S P—A 10 p c w/v solution of the pure salt, $\text{Ba}(\text{NO}_3)_2$. Barium Nitrate should answer the tests described under Barium Chloride *U S P*, but its aqueous solution acidulated with Nitric Acid should not be rendered turbid by Silver Nitrate T S, indicating the absence of Chlorides

BENZIN OR PETROLEUM BENZIN

B P—Not included

P G—The official 'Benzinum Petrolei' of the Pharmacopœia Sp gr 0.640 to 0.670 Boiling point between 50° and 75° C (122° and 167° F)

U S P—The official 'Benzinum Purificatum' of the Pharmacopœia Sp gr 0.688 to 0.660 at 25° C (77° F) Boiling point 45° to 60° C (113° to 140° F)

BENZOL

B P—The official Benzol of the Pharmacopœia

P G—A colourless liquid, Sp gr 0.880 to 0.890 Boiling point 80° to 82° C (176° to 179.6° F)

U S P—Benzene or Benzole is a colourless, transparent liquid, represented by

the chemical formula C_6H_6 . Sp gr 0.871 at $25^\circ C$ ($77^\circ F$). It congeals at $5.2^\circ C$ ($41.3^\circ F$) and boils at $80.4^\circ C$ ($176.7^\circ F$). It is insoluble in Water, but soluble in 4 parts of Alcohol and in Ether.

Concentrated Sulphuric Acid when shaken with an equal volume of Benzene should not become coloured. 2 cc of Benzene, 0.5 cc of Sulphuric Acid and 1 drop of fuming Nitric Acid shaken together should not produce a green or blue tint.

BENZOLATED AMYL ALCOHOL

B P—A mixture of Benzol and Amyl Alcohol, containing about a quarter of its volume of Amyl Alcohol, and prepared by mixing 30 parts of Benzol with 10 of Amyl Alcohol, any Water which separates out being removed by decantation. It is chiefly used as a solvent for the mixed alkaloids in the assay of Cinchona Bark and its galemical preparations.

P G
U S P } Not included.

BISMUTH OXYNITRATE

B P—The official salt of the Pharmacopœia

P G
U S P } Not included.

BORAX

B P } The official salt of the respective Pharmacopœias.
P G }
U S P —Not included

BORIC ACID, SOLUTION OF

B P—A filtered 2.5 p c w/v solution of the official acid in Alcohol (90 p c)

P G
U S P } Not included.

BROMINE

B P—Commercial Bromine

P G—The 'Bromum' of the Pharmacopœia

U S P—Not included

BROMINE, SOLUTION OF

B P—A 0.66 p c v/v solution of Bromine in Distilled Water. Should be kept protected from light.

P G—A saturated aqueous solution

U S P—A 1 p c v/v aqueous solution of the 'Bromum' of the Pharmacopœia

CADMIUM IODIDE

B P—The crystalline commercial salt represented by the chemical formula CdI_2 .

P G
U S P } Not included

CADMIUM IODIDE, SOLUTION OF

B P—A 5 p c w/v filtered aqueous solution of the above salt

P G
U S P } Not included.

CALCIUM CARBONATE

B P—Commercial White Marble or Calc Spar

P G—The salt should be free from Chlorides

U S P—Not included

CALCIUM CHLORIDE SOLUTION

B P—A filtered 10 p c w/v aqueous solution of the fused salt

P G—A 10 p c w/w aqueous solution of the crystallised salt

U S P—A 10 p c w/v aqueous solution of the crystallised salt $\text{CaCl}_2 + 6\text{H}_2\text{O}$

CALCIUM HYDROXIDE

B P—The slaked Lime of the Pharmacopœia

P G—Calcium Hydroxide

U S P—Not included

CALCIUM HYDROXIDE SOLUTION

B P }
P G } The Lime Water official in the respective Pharmacopœias
U S P }

CALCIUM SULPHATE

B P—The pure native salt represented by the chemical formula $\text{CaSO}_4, 2\text{H}_2\text{O}$

P G }
U S P } Not included

CALCIUM SULPHATE SOLUTION

B P—A saturated filtered aqueous solution of Calcium Sulphate prepared by triturating 1.25 grammes of the salt with 10 c c of Distilled Water in a porcelain mortar and adding 90 c c of Water, filtering after allowing to stand for some time

P G—A saturated aqueous solution of the salt

U S P—A saturated aqueous solution of native Gypsum, $\text{CaSO}_4, 2\text{H}_2\text{O}$, obtained by shaking the powdered crystals of Gypsum in a bottle nearly full with Water at intervals during 12 hours, then decanting the clear saturated solution when required. The solubility at 25°C (77°F) is 1 in 878

CARBON BISULPHIDE (DISULPHIDE)

B P }
U S P } The Carbon Disulphide official in the respective Pharmacopœias

P G—A colourless, volatile, neutral liquid. Boiling point 46°C (114.8°F)
 Sp gr 1.272

CHLORINATED SODA SOLUTION

B P—The official Liquor Sodæ Chlorinatæ

P G }
U S P } Not included

CHLORINE SOLUTION

B P—A freshly prepared, saturated solution of Chlorine in Water, obtained by saturating Water with the purified and washed gas obtained by the decomposition of Hydrochloric Acid and Manganese Dioxide. It possesses a gravity of 1.008, and contains about 0.5 p c of available Chlorine

U S P—The Liquor Chlori Co of the Pharmacopœia. It contains about 0.4 p c of Chlorine and should be freshly prepared when required

P G—The 'Aqua Chlorata' of the Pharmacopœia. A solution of Chlorine containing not less than 0.4 p c and not more than 0.5 p c w/w of Chlorine.

CHLOROFORM

B P }
P G } The Chloroform official in the respective Pharmacopœias
U S P }

The *U S P* requires that it should be strictly neutral to Litmus paper

CHROMIC ACID SOLUTION

B P—The Liquor Acidi Chromici of the Pharmacopœia

P G—A 3 p c w/w aqueous solution of Chromic Acid prepared when required

U S P—Not included

CITRIC ACID

B.P.—The acid official in the Pharmacopœias*P.G.*
U.S.P. } Not included

COBALTOUS NITRATE SOLUTION

B.P.
P.G. } Not included*U.S.P.*—A 10 p c w/v solution of Cobaltous Nitrate obtained by dissolving 1 commercial Cobaltous Nitrate in 10 c c of WaterCommercial crystalline Cobalt Nitrate $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, may be used in making the solution if, when dissolved in Water and the *C.* by Ammonium Sulphide T.S., the filtrate leaves no residue after gitting

COLLODION

B.P.
P.G. } The Collodion official in the respective Pharmacopœias
U.S.P.—Not included

COPPER

B.P.
U.S.P. } The metal Cu in the form of wire, foil, or turnings
P.G.—Not included

COPPER OXYACETATE

B.P.—Pure commercial Verdigris*P.G.*
U.S.P. } Not included

COPPER ACETATE SOLUTION

B.P.—A filtered 10 p c w/v aqueous solution of Copper Oxyacetate prepared by digesting a weighed quantity of 10 grammes of finely-powdered Copper Oxyacetate in a mixture of 20 c c of Acetic Acid, and 10 c c of Water at a temperature of 100°C (212°F), evaporating to dryness, digesting the resulting residue in 80 c c of boiling Water, adding sufficient Water to produce a volume of 100 c c

COPPER SULPHATE

B.P.—The salt represented by the chemical formula $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$, official in the Pharmacopœias*P.G.*
U.S.P. } Not included

COPPER SULPHATE SOLUTION

B.P. } A 10 p c w/v solution of Copper Sulphate in Water (filtered if
U.S.P. } necessary, *B.P.*)The *U.S.P.* uses the salt, $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$, official in the text*P.G.*—Not included

DIPHENYLAMINE TEST-SOLUTION

B.P.
P.G. } Not included*U.S.P.*—A 0.2 p c w/v solution of Diphenylamine in diluted Sulphuric Acid. Diphenylamine, $(\text{C}_6\text{H}_5)_2\text{NH}$, is in greyish-white or colourless crystals slightly soluble in Water, more soluble in acids. It has a peculiar, aromatic odour, and melts at 54°C (129.2°F). The T.S. should be colourless.

ETHER

$\left. \begin{array}{l} B P \\ P G \\ U S P \end{array} \right\}$ The Ether official in the respective Pharmacopœias

The *U S P* requires that it should be strictly neutral to moistened Litmus paper

FERRIC AMMONIUM SULPHATE SOLUTION

B P—Not included

P G—A solution of 1 part of Ferric Ammonium Sulphate in a mixture of Water 8 parts, and diluted Sulphuric Acid 1 part, to be prepared when required

U S P—A 10 p c w/v aqueous solution of Ferric Ammonium Sulphate $\text{FeNH}_4(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, *U S P*

FERRIC CHLORIDE

B P—The pure anhydrous commercial Ferric Chloride

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

FERRIC CHLORIDE SOLUTION

B P—A 5 p c w/v aqueous solution of the above pure anhydrous Ferric Chloride. It should be filtered if necessary

P G—Use the 'Liquor Ferri Sesquichlorati' of the Pharmacopœia diluted when necessary as directed

U S P—A 10 p c w/v solution of the Ferric Chloride of the Pharmacopœia in Distilled Water

FERRIC SULPHATE SOLUTION

B P—Use the 'Liquor Ferri Persulphatis' of the Pharmacopœia

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

FERROUS SULPHATE

B P—The salt official in the Pharmacopœia represented by the chemical formula $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

FERROUS SULPHATE SOLUTION

B P—A freshly prepared, filtered 2 p c w/v solution of Ferrous Sulphate in Distilled Water

P G—A solution of 1 part of Ferrous Sulphate in a mixture of 1 part of Water, and 1 part of diluted Sulphuric Acid to be prepared when required

U S P—The *U S P* directs that a clear crystal of Ferrous Sulphate $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ be dissolved in about 10 parts of Water which has been previously boiled to expel air. The solution should be freshly prepared immediately before use

FERROUS SULPHIDE

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Not included

U S P—A heavy solid represented by the chemical formula FeS . It is in the form of black or brownish black irregular masses, or fused into sticks, and is soluble in diluted Sulphuric Acid or diluted Hydrochloric Acid, with copious evolution of Hydrogen Sulphide

GELATIN TEST SOLUTION

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Not included

U S P—A freshly made solution of 1 gramme of the Gelatin official in the Pharmacopœia in 50 c c of Water. It is made with the aid of a gentle heat, and filtered if necessary

GLYCERIN.

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ The Glycerin official in the respective Pharmacopœias
U S P—Not included

HYDROCHLORIC ACID

$\left. \begin{array}{l} B P \\ P G \\ U S P \end{array} \right\}$ The Acidum Hydrochloricum of the respective Pharmacopœias

The *U S P* requires that the acid for use as a reagent should conform to the following additional tests. The addition of 1 c c of Barium Chloride T S to 1 c c of the acid diluted with 9 c c of Water should cause no turbidity within 24 hours. A crystal of Diphenylamine dropped into the acid should not turn blue (absence of free Chlorine).

HYDROCHLORIC ACID, DILUTED

B P—The diluted acid of the Pharmacopœia containing 10.58 p c of Hydrochloric Acid.

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

HYDROCHLORIC ACID, FUMING

$\left. \begin{array}{l} B P \\ U S P \end{array} \right\}$ Not included

P G—A colourless fuming liquid answering the tests of purity for the Hydrochloric Acid of the Pharmacopœia. Sp. gr. 1.190.

HYDROCHLORIC ACID, GASEOUS

B P—Hydrochloric Acid in the dry gaseous form, directed to be prepared from Sulphuric Acid and Sodium Chloride.

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

HYDROGEN PEROXIDE SOLUTION

B P—The official Liquor of the Pharmacopœia.

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

HYDROGEN SULPHIDE

$\left. \begin{array}{l} B P \\ U S P \end{array} \right\}$ The washed gas represented by the chemical formula H_2S obtained by the action of Hydrochloric Acid on Ferrous Sulphide. The *B P* employs Hydrochloric Acid, the *U S P* diluted Sulphuric Acid.

P G—Not included.

The *B P* states that the gas may be used after having been passed through two wash bottles containing Water, and the *U S P* requires that after generation the gas be treated as described under Hydrogen Sulphide Solution.

HYDROGEN SULPHIDE SOLUTION

B P—An aqueous solution of Hydrogen Sulphide gas. It should have a strong odour of Hydrogen Sulphide and should give a copious black precipitate with Lead Subacetate T S.

P G—A saturated aqueous solution of Hydrogen Sulphide.

U S P—A saturated aqueous solution of Hydrogen Sulphide. It directs that 1000 c c of the solution be prepared by treating 20 grammes of Ferrous Sulphide with 20 c c of Sulphuric Acid (*U S P*) mixed with 250 c c of Water. The generated gas is first passed through a drying tube containing granulated Calcium Chloride, then from this through a tube of about 5 mm diameter and about 40 cm in length, containing about 5 grammes of coarsely pulverised Iodine mixed with glass wool, and finally through a wash-bottle containing a little Potassium Iodide T S. The purified gas is then passed through 1000 c c of Water contained in a bottle of 1500 c c capacity, shaking the bottle occasionally.

After absorption ceases the solution is transferred to small dark amber coloured bottles, allowing a stream of the gas to pass through each before stoppering. The bottles should be filled nearly to the top and preserved in a cool and dark place. The solution should only be used if it retains a strong odour of the gas and yields a copious precipitate of Sulphur when it is added to an equal volume of Ferric Chloride T S.

ISINGLASS

BP—A shredded gelatinous substance prepared from the sounds or swimming bladders of different species of *Acipenser*, Linn. See also p. 655

PG }
USP } Not included

ISINGLASS, SOLUTION OF

BP—The freshly prepared, filtered, aqueous 2 p.c. w/v solution of Isinglass prepared by digesting the Isinglass in Water for half an hour at a water bath temperature. The product is filtered through tow.

LEAD ACETATE

BP—The salt official in the Pharmacopœia

PG }
USP } Not included

LEAD ACETATE SOLUTION

BP }
USP } A 10 p.c. w/v solution of the salt in Distilled Water

PG—A 10 p.c. w/w solution of the salt in Distilled Water

The *BP* directs that recently boiled Distilled Water should be used and that the solution be filtered. The *USP* directs that only transparent crystals of the salt be used free from adhering Carbonate.

LEAD PEROXIDE

BP—The pure commercial Lead Peroxide represented by the chemical formula PbO_2 .

PG }
USP } Not included

LEAD SUBACETATE SOLUTION

BP }
PG } The 'Liquor Plumbi Subacetatis Fortis' official in the respective
USP } Pharmacopœias

This is the Basic Lead Acetate Test Solution of the *USP*.

LIME, SOLUTION OF CHLORINATED

BP }
USP } Not included

PG—Triturate 1 part of Chlorinated Lime with 9 parts of Water, and filter the solution. It should be prepared when required.

MAGNESIUM AMMONIO-SULPHATE SOLUTION (MAGNESIA MIXTURE, *USP*)

BP } Magnesium Sulphate, 10 grammes, Ammonium Chloride,
USP } 20 grammes, dissolve in 80 c.c. of Water and add 42 c.c. of Ammonia Solution (Aqua Ammonia, *USP*)

The *BP* directs that the mixture be allowed to stand for a few days in a well-stoppered bottle, then decanted and filtered. The *USP* states that if not perfectly clear the mixture should be filtered before using.

MAGNESIUM SULPHATE SOLUTION

BP }
USP } A 10 p.c. w/v (filtered *BP*) solution of the official salt in Distilled Water

PG—A 10 p.c. w/w solution of the salt in Water

MANGANESE PEROXIDE

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Powdered native Pyrolusite, MnO_2
U.S.P.—Not included

MERCURIC CHLORIDE SOLUTION

$\left. \begin{array}{l} B P \\ U S P \end{array} \right\}$ A 5 p c w/v filtered aqueous solution of the official salt
P.G.—A 5 p c w/w aqueous solution of the official salt

MERCURIC NITRATE SOLUTION

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Not included
U.S.P.—The 'Liquor Hydrargyri Nitratis' of the Pharmacopœia

MERCUROUS NITRATE SOLUTION

B.P.—A solution of Mercurous Nitrate obtained by dissolving 1 gramme of Mercury in a mixture of 0.5 c c of Water and 0.5 c c of Nitric Acid, the mixture being allowed to remain at rest for 24 hours in a cool dark place and the crystalline residue dissolved in 100 c c of Water

U.S.P.—A solution of Mercurous Nitrate prepared by mixing 10 grammes of Mercury, 5 c c of pure Nitric Acid, and 5 c c of Distilled Water, and setting aside in a cool dark place. The crystals formed during 24 hours are, after draining, dissolved in 100 c c of Water. The solution should be preserved in a dark amber-coloured bottle into which a small quantity of Mercury has been introduced

P.G.—Not included

MERCURY

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Not included

U.S.P.—Rectified purified wood Alcohol CH_3OH Sp gr, about 0.812 at $25^\circ C$ ($77^\circ F$). It should be free from pyroligneous odour

MICROCOSMIC SALT

B.P.—The commercial salt represented by the chemical formula $NaNH_2HPO_4 \cdot 4H_2O$.

$\left. \begin{array}{l} P G \\ U S P \end{array} \right\}$ Not included

NAPHTHYLAMINE ACETATE SOLUTION

$\left. \begin{array}{l} B P \\ P G \end{array} \right\}$ Not included

U.S.P.—Boil 0.1 gramme of Alphanaphthylamine Acetate ($C_{10}H_7NH_2 \cdot HC_2H_3O_2$) in 20 c c of Distilled Water, filter through cotton, and mix the filtrate with 180 c c of diluted Acetic Acid (10 p c of absolute Acid). Only freshly Distilled Water should be employed in preparing this reagent, which must be kept in well-corked bottles in the light

NITRIC ACID

$\left. \begin{array}{l} B P \\ P G \\ U S P \end{array} \right\}$ The official Nitric Acid of the respective Pharmacopœias

NITRIC ACID, CRUDE

$\left. \begin{array}{l} B P \\ U S P \end{array} \right\}$ Not included

P.G.—A clear colourless or yellowish coloured liquid, fuming on heating, completely volatile on warming. Sp gr 1.390 to 1.400. It contains a trace of pure acid.

NITRIC ACID, DILUTED

B P—The official diluted acid of the Pharmacopœia

P G—Nitric Acid 1 part, Water 1 part, prepared when required

U S P—Not included

NITRIC ACID, FUMING

B P—Nitric Acid having a sp gr of 1.5

P G—A clear reddish brown liquid, completely volatile on warming. It evolves suffocating yellowish red fumes. Sp gr 1.486 to 1.500. It contains 86 p c of pure acid.

U S P—The commercial red fuming acid. Sp gr 1.437 at 25° C (77° F). It should be carefully kept in glass stoppered bottles in a cool place.

OLIVE OIL

B P—The Olive Oil of the Pharmacopœia

P G } Not included

U S P }

OXALIC ACID

B P—Not included

P G—The air dried acid. It should leave no residue when ignited on platinum foil.

U S P—Pure Oxalic Acid represented by the chemical formula $\text{H}_2\text{C}_2\text{O}_4 + 2\text{H}_2\text{O}$, 10 grammes on ignition on platinum foil should leave no residue. It should be completely soluble in 12 parts of Water at 25° C (77° F). For the preparation of test and volumetric solutions, commercial Oxalic Acid should be purified as follows: To 1 part of the Acid add 10 parts of cold Water, and shake until the latter is saturated. Filter off the solution from the undissolved crystals, evaporate the filtrate to about three fourths of its volume, and set it aside so that the fixed salts which it contains may crystallise out. Carefully decant the liquid from the crystals, concentrate it by evaporation, and set it aside to crystallise, stirring occasionally to prevent the formation of large crystals, which might enclose moisture. Drain the crystals in a funnel, dry them carefully on blotting paper, and preserve them in well stoppered bottles.

OXALIC ACID SOLUTION

B P } Not included

P G }

U S P—A 10 p c w/v aqueous solution of pure Oxalic Acid

PALLADOUS CHLORIDE SOLUTION

B P } Not included

P G }

U S P—A 5 p c w/v aqueous solution of Palladous Chloride PdCl_2 . The solution should be preserved in a glass stoppered bottle.

PHENOL

B P—The 'Acidum Carbolicum' of the Pharmacopœia

P G } Not included in the reagent list

U S P }

PHENOL SOLUTION

B P } Not included

U S P }

P G—A 5 p c w/w aqueous solution of Carbolic Acid

PICRIC ACID

B P—Trinitrophenol represented by the formula $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH}$. See also Acidum Picricum

P G } Not included

U.S.P }

PICRIC ACID SOLUTION

B.P.—A 0.6 p.c. w/v aqueous solution of Picric Acid

P.G.—Not included

U.S.P.—A 1 p.c. w/v aqueous solution of pure, distinctly crystalline Picric Acid $C_6H_2(NO_2)_3OH$. Cool the solution and filter if necessary

PLATINIC CHLORIDE SOLUTION

B.P.—A 1 p.c. solution of Platinum Tetrachloride, equivalent to 13.2 p.c. w/v Chloro-platinum Acid, obtained by heating 5 grammes of commercial platinum foil to a temperature of about $80^\circ C$ ($176^\circ F$) with 30 c.c. of Hydrochloric Acid, and very gradually adding 5 c.c. of Nitric Acid. The liquid is evaporated to dryness on a water bath moistened with Hydrochloric Acid, and again evaporated and the residue dissolved in sufficient Water to measure 100 c.c.

P.G.—A 5 p.c. w/w solution of Chloro-platinum Acid in Water

U.S.P.—A solution of 2.6 grammes of Chloro-platinum Acid $H_2PtCl_6 \cdot 6H_2O$ in 20 c.c. of Water. It is required that if a small portion of this solution be evaporated to dryness and the residue ignited, pure metallic Platinum should remain, which should yield nothing soluble in Nitric Acid

POTASSIO-MERCURIC IODIDE ALKALINE SOLUTION (NESSLER'S)

B.P.—A solution prepared by dissolving a mixture of 3 grammes of Potassium Iodide and 1.25 grammes of Mercuric Chloride in 80 c.c. of Water. A cold saturated aqueous Mercuric Chloride Solution drop by drop with constant agitation until a faint permanent red precipitate ensues, adding 12 grammes of Sodium Hydroxide, 1 or 2 drops of the cold saturated aqueous Mercuric Chloride Solution and diluting with Water to measure 100 c.c.

U.S.P.—To a solution of 5 grammes of Potassium Iodide in 5 c.c. of Water gradually add in portions a saturated aqueous solution of Mercuric Chloride with constant agitation until a slightly red precipitate remains undissolved. Add 15 grammes of Potassium Hydroxide, and, when this has dissolved, 0.5 c.c. more of the saturated aqueous solution of Mercuric Chloride, then dilute with Water to 100 c.c. Allow the precipitate to subside and draw off the clear fluid. 2 c.c. of this reagent when added to 50 c.c. of Water containing 0.05 milligramme of Ammonia should produce at once a yellowish-brown coloration

P.G.—Not included

POTASSIO-MERCURIC IODIDE SOLUTION MAYER'S REAGENT

B.P. } Not included

P.G. }

U.S.P.—Dissolve 1.344 grammes of Mercuric Chloride in 60 c.c. of Water and 5 grammes of Potassium Iodide in 10 c.c. of Water. Mix the two solutions, and then add sufficient Water to make the mixture measure 100 c.c.

POTASSIUM ACETATE SOLUTION

B.P.—A 10 p.c. w/v filtered aqueous solution of Potassium Acetate

P.G.—The Liquor Kali Acetici of the Pharmacopœia prepared by gradually adding 24 parts of Potassium Bicarbonate to 50 parts by weight of diluted Acetic Acid, *P.G.*, heating the solution to boiling point then neutralising with Potassium Bicarbonate and diluting the cooled liquid with Water until the solution has a sp. gr. of 1.176 to 1.180

U.S.P.—Not included

POTASSIUM ACID TARTRATE SOLUTION

B.P.—A saturated solution of Potassium acid salt obtained by dissolving it in Distilled Water and filtering

P.G. } Not included.

U.S.P. }

POTASSIUM BICHROMATE

BP—The official salt of the Pharmacopœia See Potassu Bichromas

PG—Not included

USP—Pure Potassium Dichromate The pure salt $K_2Cr_2O_7$, answering the following tests of purity in addition to the official requirements. In a solution of 0.5 gramme of the salt in 10 c.c. of Water rendered acid by 0.5 c.c. of Nitric Acid, no turbidity should be produced by Barium Chloride T.S. (absence of Sulphates). 10 c.c. of an aqueous solution of the salt (1-20) should give no turbidity with 1 c.c. of Ammonia Water followed by 1 c.c. of Ammonium Oxalate T.S. (absence of Calcium). If to a solution of 0.5 gramme of the salt in 20 c.c. of Water sufficient Sulphurous Acid be added to impart a strong odour of the reagent and the mixture be boiled for about 3 minutes and cooled, the addition of 1 c.c. of Nitric Acid and a few drops of Silver Nitrate V.S. should produce no turbidity (absence of Chlorides). Should the official salt not answer these tests it may be purified by repeated recrystallisation until it does so. A hot saturated aqueous solution of the salt is rapidly cooled with agitation and the granular crystals collected on a plum filter, washed with cold Water to remove the mother liquor, drained and then dried at $120^\circ C$ ($245^\circ F$). This recrystallisation is repeated as often as is necessary.

POTASSIUM BICHROMATE SOLUTION

BP—Not included

PG—A 5 p.c. w/w aqueous solution of the official salt

USP—A 10 p.c. w/v aqueous solution of the pure salt

POTASSIUM BROMATE

BP } Not included
PG }

USP—The salt represented by the chemical formula $KBrO_3$. It occurs as white cubical crystals or a granular crystalline powder, and has a pungent saline taste. It should respond to the following tests.—**Solubility** 1 in 15.5 of Water at $25^\circ C$ ($77^\circ F$), 1 in 2 of boiling Water, slightly in Alcohol. **Litmus** An aqueous solution should be neutral. **Sulphuric Acid** The aqueous solution of the salt should not at once yield a yellow colour on the addition of the diluted acid, but Sulphuric Acid added to the salt causes decomposition with evolution of Bromine. **Nitric Acid** Also causes decomposition with evolution of Bromine. **Heat** At $350^\circ C$ ($662^\circ F$) decomposition occurs and Oxygen is given off. **Volumetric determination** Dissolve 0.1 gramme of the salt which has been dried at $100^\circ C$ ($212^\circ F$) and 2 grammes of Potassium Iodide in 25 c.c. of Water contained in a glass stoppered bottle of about 100 c.c. capacity, then add 5 c.c. of Hydrochloric Acid, well stopper the bottle and set aside for ten minutes. On titrating this mixture with Tenth normal Sodium Thiosulphate V.S. not less than 36.1 c.c. of the V.S. should be required to discharge the colour, corresponding to 99.8 p.c. of pure salt.

Note—Potassium Bromate should not be titrated or heated with organic or easily oxidisable substances.

POTASSIUM CARBONATE SOLUTION

BP—A 10 p.c. w/v filtered aqueous solution of Potassium Carbonate

PG—The 'Liquor Kalii Carbonici' of the Pharmacopœia made by dissolving 11 parts of the salt in 20 parts of Water, filtering the solution and diluting with Water if necessary to a sp. gr. of 1.330 to 1.334. It is an aqueous solution containing 33.3 p.c. w/w of Potassium Carbonate.

USP—A 10 p.c. w/v aqueous solution of anhydrous Potassium Carbonate. The anhydrous salt is prepared by heating the official salt to $130^\circ C$ ($266^\circ F$).

POTASSIUM CHLORATE

BP—The official salt of the Pharmacopœia See Potassu Chloras

PG } Not included
USP }

POTASSIUM CHROMATE

BP—The pure neutral commercial salt in yellow crystals represented by the chemical formula K_2CrO_4 ,

PG }
USP } Not included

POTASSIUM CHROMATE SOLUTION

BP—A 10 p c w/v in

solution of Potassium Chromate

PG—A 5 p c w/v of yellow Potassium Chromate free from Chlorine

USP—A 10 p c w/v aqueous solution of yellow Potassium Chromate. The red by the addition of Silver Nitrate T.S. to a few drops of the solution in a little Distilled Water, should be entirely soluble in Nitric Acid (absence of Chlorides). It should be free from Sulphates, equal volumes of the solution and diluted Hydrochloric Acid yielding no precipitate with Barium Chloride T.S. Another portion of the solution should give no turbidity with Ammonia Water or Ammonium Oxalate T.S. (absence of alkali earths). A solution of 0.1 gramme of the salt in 20 c c of Water should not become red on the addition of a few drops of Phenolphthalein T.S. (limit of free alkalis)

POTASSIUM CYANIDE

BP—The Potassium Cyanide, KCN, of commerce yielding not less than 90 p c of pure salt

PG }
USP } Not included

POTASSIUM CYANIDE SOLUTION

BP }
USP } A 10 p c w/v aqueous solution of Potassium Cyanide (*BP* filtered)

The *USP* uses the official salt of the Pharmacopoeia and directs that the solution be prepared when required

PG—Not included

POTASSIUM FERRICYANIDE

BP—The salt represented by the chemical formula $K_3FeC_6N_6$. It occurs as red crystals and should be free from Ferrous salt as ascertained by its aqueous solution failing to give a precipitate or blue coloration with a dilute solution of a pure Ferric salt

PG }
USP } Not included

POTASSIUM FERRICYANIDE SOLUTION

BP—A freshly prepared 5 p c w/v aqueous filtered solution of crystallised Potassium Ferricyanide

PG—A freshly prepared 5 p c w/v aqueous solution of Potassium Ferricyanide in crystals which have been previously washed with Water

USP—A solution of 1 part of Potassium Ferricyanide in about 10 parts of Water, freshly made when required. The absence of Ferrous salt is proved by its failure to give any turbidity or shade of green with Ferric Chloride T.S. when diluted with Water, only a brown tint can be detected. Potassium Ferricyanide should be free from Sulphates and Chloride

POTASSIUM FERRIOXYANIDE

BP—The salt in yellow crystals obtained by the fusion of a mixture of Potassium Carbonate, Iron and nitrogenous organic matter. It is represented by the chemical formula $K_4FeC_6N_8, 3H_2O$

PG—Not included

USP—The yellow crystalline salt represented by the formula $K_4Fe(CN)_6, 3H_2O$.

POTASSIUM FERROCYANIDE SOLUTION

BP—A 5 p c w/v aqueous filtered solution of Potassium Ferrocyanide in crystals

PG—A freshly prepared 5 p c w/w aqueous solution of Potassium Ferrocyanide

USP—A 10 p c w/v aqueous solution of the salt

POTASSIUM HYDROGEN SULPHITE

BP—Acid Potassium Sulphite of commerce represented by the chemical formula KHSO_3

PG
USP } Not included

POTASSIUM HYDROXIDE

BP—The Potassa Caustica of the Pharmacopœia

PG
USP } Not included

POTASSIUM HYDROXIDE SOLUTION

BP
PG
USP } The solutions official in the respective Pharmacopœias

POTASSIUM HYDROXIDE SOLUTION (ALCOHOLIC)

BP—A 10 p c w/v filtered solution of Potassium Hydroxide in Alcohol (90 p c)

PG—A 10 p c w/w solution of fused Potassium Hydroxide in Alcohol (90 to 91 2 p c)

USP—Use the Half normal Alcoholic Potassium Hydroxide V S, *q v p* 1001

POTASSIUM IODIDE

BP—The official salt of the Pharmacopœia

PG
USP } Not included

POTASSIUM IODIDE SOLUTION

BP—A 10 p c w/v filtered aqueous solution of the official Potassium Iodide

PG—A 10 p c w/w aqueous solution of the official salt

USP—A 20 p c w/v aqueous solution of the official salt. It should be kept in dark amber coloured, well stoppered bottles, and should be frequently renewed

POTASSIUM NITRATE

BP
PG } Not included

USP—The dry salt, KNO_3 , official in the Pharmacopœia. It should also be free from Chlorides and Sulphates

POTASSIUM PERMANGANATE

BP—The official salt of the Pharmacopœia

PG
USP } Not included

POTASSIUM PERMANGANATE SOLUTION

BP—The 'Liquor Potassii Permanganatis' of the Pharmacopœia

PG—A 0.1 p c w/w aqueous solution of the official salt

USP—Use the Tenth-normal Volumetric Solution

POTASSIUM SULPHATE

B P — The official salt of the Pharmacopœia
P G } Not included
U S P }

POTASSIUM SULPHATE SOLUTION

$\left. \begin{matrix} BP \\ PG \end{matrix} \right\}$ Not included
USP—A 1 p.c. w/v aqueous solution of Potassium Sulphate

POTASSIUM SULPHOCYANATE

$\left. \begin{array}{l} BP \\ PG \end{array} \right\}$ Not included

USP—The salt represented by the chemical formula $KSCN$. It is in the form of colourless prismatic crystals, which are hygroscopic in moist air. It has a cooling saline taste. **Solubility** at $25^{\circ}C$ ($77^{\circ}F$) 1 in less than 1 of Water, 1 in 10 of Absolute Alcohol. An aqueous solution of the salt (1:20) should not become turbid within 5 minutes upon the addition of Barium Chloride T.S. (limit of Sulphate). The aqueous solution (1 in 20) after the addition of 1 c.c. of diluted Hydrochloric Acid, should remain colourless (absence of Iron), and should not respond to the time-limit test for heavy metals.

POTASSIUM SULPHATE SOLUTION

$\left. \begin{matrix} B P \\ P G \end{matrix} \right\}$ Not included
 $U S P$ — Use the Tenth-normal V S

PYROXYLIN SOLUTION

B P—The Collodion official in the Pharmacopœia
P G } Not included
U S P }

SILVER AMMONIO-NITRATE SOLUTION

BP—A solution obtained by dissolving 2.5 grammes of Silver Nitrate in 80 c.c. of Distilled Water, and adding Ammonia Solution drop by drop until the precipitate almost redissolves, the clear liquid is decanted from the precipitate, and sufficient Water added to measure 100 c.c.

U S P—Prepared by adding, drop by drop, Ammonia Water *qs* to a solution of 1 gramme of Silver Nitrate in 20 cc of Water until solution of the precipitate formed is nearly, but not entirely, effected. Filter and preserve the solution in dark amber-coloured and well-stoppered bottles.

SILVER NITRATE SOLUTION

USP—For ordinary purposes use the Tenth normal Volumetric Solution

SILVER SULPHATE SOLUTION

$\frac{BP}{PG}$ } Not included

USP—To a solution of 1 gramme of the official Silver Nitrate in 0.5 cc of warm Water add 1.5 cc of pure concentrated Sulphuric Acid, cool the solution, and pour off the acid liquid. The residue is washed with Silver Solution, filtered, and poured into a bottle, and the residue is washed with Water. Decant the supernatant quantity of this saturated solution and repeat.

SODIUM ACETATE

B P—The salt represented by the chemical formula CH_3COONa , $3\text{H}_2\text{O}$. The use of the pure commercial salt is officially permitted.

P G
U S P } Not included

SODIUM ACETATE SOLUTION

B P—A 10 p c w/v filtered aqueous solution of the salt.

P G—A 20 p c w/w aqueous solution of the official salt.

U S P—A 10 p c w/v aqueous solution of the official salt.

SODIUM ARSINATE

B P—The official salt of the Pharmacopœia.

P G
U S P } Not included

SODIUM BICARBONATE

B P—The official salt of the Pharmacopœia.

P G
U S P } Not included

SODIUM BICARBONATE SOLUTION

B P
U S P } Not included

P G—Dissolve 1 part of the powdered salt in 19 parts of Water with gentle stirring.

SODIUM BITARTRATE SOLUTION

B P
P G } Not included

U S P—To a solution of 3.5 grammes of the official Tartaric Acid in about 80 c c of boiling Water, add gradually, in small portions, Monohydrated Sodium Carbonate $\text{Na}_2\text{CO}_3 \cdot \text{H}_2\text{O}$ until the solution has a neutral reaction, to this liquid is now added 3.5 grammes of Tartaric Acid, and after filtering and cooling sufficient Water is added to the solution to measure 100 c c. This solution should be freshly prepared when required.

SODIUM CARBONATE

B P—The official salt of the Pharmacopœia.

P G—Not included.

U S P—The official monohydrated salt $\text{Na}_2\text{CO}_3 \cdot \text{H}_2\text{O}$. It should respond to the tests of the Pharmacopœia, and should be absolutely free from Chloride and Sulphate.

SODIUM CARBONATE SOLUTION

B P—A 10 p c w/v filtered aqueous solution of the official salt.

P G—A 20 p c w/w aqueous solution of the official salt.

U S P—A 10 p c w/v aqueous solution of the 'Sodium Carbonate Monohydratus' of the Pharmacopœia.

SODIUM CHLORIDE

B P—The official salt of the Pharmacopœia.

P G
U S P } Not included

SODIUM COBALTIC NITRITE SOLUTION

B P
P G } Not included

U S P—A solution of Sodium Cobaltic Nitrite $\text{Co}(\text{NO}_2)_2 \cdot 6\text{NaNO}_2 \cdot \text{H}_2\text{O}$ made by dissolving 4 grammes of Cobaltous Nitrate $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 10 grammes of Sodium Nitrite NaNO_2 in about 50 c c of Water, then adding 2 c c of Acetic

Acid (*U.S.P.*) and diluting with Water to 100 c.c. A few drops of Acetic Acid should be added to the solution from time to time, and it should not be kept longer than three months.

SODIUM HYDROGEN SULPHITE

B.P.—The salt NaHSO_3 found in commerce

P.G. }
U.S.P. } Not included

SODIUM BISULPHITE SOLUTION

B.P. }
U.S.P. } Not included

P.G.—It contains about 30 p.c. w/w of Sodium Bisulphite

SODIUM HYDROXIDE

B.P.—For the official varieties see Soda Caustica

P.G.—The *P.G.* uses the fused Caustic Soda and requires that an aqueous solution (1-6) should respond to the tests of purity given for the 'Liquor Natri Caustici' of the Pharmacopœia

U.S.P.—Not included

SODIUM HYDROXIDE SOLUTION

B.P.—A 20 p.c. w/v filtered aqueous solution of the 'Purified Sodium Hydroxide'

P.G.—The 'Liquor Natri Caustici' of the Pharmacopœia Sp. gr. 1.168 to 1.172. It contains about 15 p.c. w/w of Sodium Hydroxide

U.S.P.—The 'Liquor Sodii Hydroxidi' of the Pharmacopœia Sp. gr. 1.056 at 25° C (77° F). It contains about 5 p.c. w/w of Sodium Hydroxide

SODIUM NITRITE

B.P.—The official salt of the Pharmacopœia

P.G.—Not included

U.S.P.—The purest commercial salt, either granulated or in the form of sticks may be employed

SODIUM NITRO-PRUSSIDE SOLUTION

B.P. }
P.G. } Not included

U.S.P.—A 10 p.c. w/v solution of Sodium Nitro-prusside $\text{NaFe}(\text{NO})(\text{CN})_5$, $2\text{H}_2\text{O}$, before use by dissolving 1 part of the salt in 19 parts of Water

SODIUM PHOSPHATE SOLUTION

B.P. } A 10 p.c. w/v filtered aqueous solution of the salt official in the
U.S.P. } respective Pharmacopœias

P.G.—A 5 p.c. w/v aqueous solution of the official salt

SODIUM POTASSIUM TARTRATE

B.P.—The 'Soda Tartarata' of the Pharmacopœia

P.G. }
U.S.P. } Not included

SODIUM SULPHATE

B.P.—The official salt of the Pharmacopœia

P.G. }
U.S.P. } Not included

SODIUM SULPHATE SOLUTION

B.P.—A 10 p.c. w/v filtered aqueous solution of the official salt,

P.G. }
U.S.P. } Not included,

SODIUM SULPHITE

B P —The official salt of the Pharmacopoeia

P G } Not included
U S P }

SODIUM SULPHITE SOLUTION

B P } Not included
U S P }

P G —A 10 p.c. w/w aqueous solution of Sodium Sulphite. It should be prepared when required.

SODIUM TARTRATE SOLUTION

B P } Not included
P G }

U S P —A solution of Sodium Tartrate $\text{Na}_2\text{C}_4\text{H}_4\text{O}_6 \cdot 2\text{H}_2\text{O}$ prepared by adding gradually, in small portion, Monohydrated Sodium Carbonate *U S P* to a solution of 6.5 grammes of Tartaric Acid *U S P* in about 50 c.c. of boiling Water, until the solution has a neutral reaction. It is then filtered, cooled and made up to 100 c.c. with Water. It should be freshly prepared when required.

SODIUM THIOSULPHATE

B P —*Syn.* Sodium Hyposulphite. A crystalline salt represented by the chemical formula $\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$

P G } Not included
U S P }

SODIUM THIOSULPHATE SOLUTION

B P } Not included
P G }

U S P —Use the Tenth normal V.S.

STANNOUS CHLORIDE SOLUTION

B P —A solution of Stannous Chloride obtained by heating 20 grammes of granulated Tin with a mixture of 60 c.c. of Hydrochloric Acid and 20 c.c. of Water until the evolution of gas ceases, the undissolved Tin is allowed to remain in the liquid, to which sufficient Water should be added to measure 100 c.c.

P G —Let 5 parts of Stannous Chloride in crystals be mixed to a paste with 1 part of Hydrochloric Acid and the mixture completely saturated with dry Hydrochloric Acid gas. The solution so obtained is poured off after being allowed to deposit, and filtered through asbestos. It is a pale yellow, strongly fuming, refractive liquid with a sp. gr. of not less than 1.900. The solution mixed with 10 volumes of Spirit should not become turbid even in the course of an hour. No turbidity should be produced by Ferrum Chloride Solution (1-20) in Stannous Chloride Solution diluted with 10 volumes of Water, even after ten minutes. The solution should be preserved in small well stoppered bottles as full as possible.

U S P —A solution of 1 part Stannous Chloride in crystals, in 10 parts of Water. The crystals of Stannous Chloride are prepared by heating Tin (in foil or granules) with concentrated Hydrochloric Acid, keeping the metal in excess. When the Acid is saturated, crystals of Stannous Chloride, $\text{SnCl}_2 \cdot 2\text{H}_2\text{O}$ begin to form. These are removed and drained, and are then used in making the solution.

The solution should be preserved in well stoppered bottles containing a fragment of pure Tin or a piece of Tin foil.

For Bettendorfs test pure concentrated Hydrochloric Acid (which responds to the *U S P* tests of purity) is saturated with the freshly prepared crystals.

SULPHANILIC ACID SOLUTION

B P } Not included
P G }

U S P —A solution of 0.5 gramme of Sulphanilic Acid $\text{C}_6\text{H}_4(\text{NH}_2)(\text{SO}_3\text{H})$ (Para amidobenzenesulphonic acid) in 150 c.c. of diluted Acetic Acid (10 p.c.

absolute Acetic Acid) Only freshly distilled Water should be employed in preparing the diluted Acetic Acid and the reagent should be kept in well-stoppered bottles

SULPHUR

BP—The 'Sulphur Sublimatum' of the Pharmacopœia

PG } Not included
USP }

SULPHURIC ACID

BP } The acids official in the respective Pharmacopœias
PG }

USP—The official acid may be used as a reagent for most purposes, provided it is of the required degree of purity. If 'concentrated' Sulphuric Acid be specially directed in a test the strongest acid, specific gravity not less than 1.84 at 25°C (77°F), should be used. This acid should respond to the official tests of purity and conform to the following additional test—If 1 c.c. of Diphenylamine dissolved in 10 c.c. of Sulphuric Acid be carefully poured, as a separate layer, upon 5 c.c. of Sulphuric Acid contained in a test-tube, no distinct blue colour should appear in the zone of contact.

SULPHURIC ACID, DILUTED

BP } The diluted acid of the respective Pharmacopœias
PG }

USP—Not included

SULPHUROUS ACID (SOLUTION)

BP—The 'Acidum Sulphurosum' of the Pharmacopœia

PG—The reagent is prepared when required by acidulating a freshly prepared solution (1-10) of Sodium Sulphite with diluted Sulphuric Acid

USP—Not included

TALC (POWDERED)

BP—A purified Magnesium Silicate *See* Talcum

PG } Not included
USP }

TANNIC ACID SOLUTION

BP—A freshly prepared 10 p.c. w/v aqueous solution of Tannic Acid

PG—A 5 p.c. w/v solution of Tannic Acid in Water, prepared when required

USP—A solution of 1 gramme of Tannic Acid in 1 c.c. of Alcohol diluted with Water to 10 c.c.

TARTARIC ANTIMONY SOLUTION

BP—A freshly prepared 5 p.c. w/v filtered solution of Tartarated Antimony in boiling Water

PG } Not included.
USP }

TARTARIC ACID

BP—The official 'Acidum Tartaricum' of the Pharmacopœia.

PG } Not included
USP }

TARTARIC ACID SOLUTION.

BP—A solution obtained by dissolving 12.5 grammes of Tartaric Acid in 65 c.c. of Water, mixing with 25 c.c. of Alcohol (90 p.c.) and adding sufficient Water to measure 100 c.c.

PG—A freshly prepared 20 p.c. w/v aqueous solution of Tartaric Acid

USP—A freshly prepared solution of 1 part of the Acidum Tartaricum in 3 parts of Water.

TIN

B P } The metal Tin in the granulated form. The *B P* requires that it
U S P } should not respond to the tests for Lead, Copper, Iron or Zinc.
The *U S P* requires that it should be free from Lead, indicated by its solution in Hydrochloric Acid failing to give a precipitate with Potassium Sulphate T.S. When tested by the modified Gutzeit's test replacing the Zinc by Tin, the diluted Hydrochloric Acid by Hydrochloric Acid *U S P*, and adding 1 drop of Platinic Chloride T.S., the Mercuric Chloride cap should not become coloured within the time required for the solution of the metal (absence of Arsenic).

P G - Tin foil free from Lead is to be employed.

TURPENTINE, OIL OF

B P -The 'Oleum Terebinthinae' of the Pharmacopœia

P G }
U S P } Not included

URANIUM NITRATE

B P -Pure commercial Uranium Nitrate in crystals

P G }
U S P } Not included

URANIUM NITRATE SOLUTION

B P -A 5 p c w/v aqueous solution of Uranium Nitrate

P G }
U S P } Not included

WATER

B P -The 'Aqua Destillata' of the Pharmacopœia

P G }
U S P } Not included

ZINC

B P -The metal in sheet or granular form. See Zincum

P G -The *P G* employs also Zinc filings

U S P -The pure metallic Zinc of the Pharmacopœia

VOLUMETRIC ANALYSIS.

STANDARDISED SOLUTIONS EMPLOYED IN VOLUMETRIC ANALYSIS

AND

INDICES OF NEUTRALITY

SOLUTIONS standardised to contain definite molecular equivalents of various chemical substances are used in the British, United States, and German as well as other countries for the purpose of determining the quantity of other substances which enter into reaction in accordance with the laws of chemical equivalence and which are present in unknown quantity.

A Normal Solution is defined as one containing in each 1000 cc. such an amount of the active constituent as will combine with, replace or oxidise 1 gramme of Hydrogen. The *BP* does not define what is meant by a Normal Solution. The *USP* defines Normal Volumetric Solutions as those which contain in 1 litre in any stated reaction the chemical equivalent of 1 gramme of Hydrogen. It draws attention to the relative weight in grammes required for 1 litre, depending on whether the molecule of the active ingredient is univalent, divalent or trivalent. The *BP* employs measuring apparatus which is adjusted at a temperature of 15.5° C (60° F). The *USP* mentions that it is absolutely necessary that the solutions should be amongst themselves accurately in their graduation at 15.5° C (60° F) but that it is immaterial what standard temperature has been selected for the graduation of the vessels. All the *USP* Volumetric Solutions must be prepared at a temperature of 25° C (77° F), and it is further required that in carrying out titrations with these solutions the temperature should not be below 21° C (69.8° F) nor above 29° C (84.2° F). The *PG* gives no general directions for the temperatures to be observed nor the method to be followed in preparing solutions for Volumetric analysis.

The following list below shows the Volumetric Solutions which are official in the three Pharmacopœias dealt with in this volume, *eg*, the *BP*, *USP* and *PG*.

BP Volumetric Sulphuric Acid Solution

Deci-normal Volumetric Sulphuric Acid Solution

Volumetric Sodium Hydroxide Solution

Deci-normal Volumetric Sodium Hydroxide Solution

Normal Alcoholic Sodium Hydroxide Solution

Deci-normal Alcoholic Sodium Hydroxide Solution

Normal Alcoholic Potassium Hydroxide Solution

Deci-normal " " Potassium Hydroxide Solution.

Volumetric Iron

Volumetric Potassium Bichromate Solution

Volumetric Silver Nitrate Solution

Volumetric Sodium Thio-sulphate Solution

USP Normal Volumetric Sulphuric Acid Solution

Half-normal Volumetric Sulphuric Acid Solution

Tenth-normal Volumetric Sulphuric Acid Solution

Fiftieth-normal Volumetric Sulphuric Acid Solution

Double normal Volumetric Sodium Hydroxide Solution

Normal " " Sodium Hydroxide Solution

Normal " " Potassium Hydroxide Solution

Tenth normal Volumetric Potassium Hydroxide Solution
 Fifteenth normal Volumetric Potassium Hydroxide Solution
 Hundredth-normal Volumetric Potassium Hydroxide Solution
 Half normal Volumetric Alcoholic Potassium Hydroxide Solution
 Normal Volumetric Hydrochloric Acid Solution
 Half normal Volumetric Hydrochloric Acid Solution
 Tenth-normal Volumetric Oxalic Acid Solution
 Tenth-normal Volumetric Iodine Solution
 Tenth normal Volumetric Bromine Solution
 Tenth normal Volumetric Potassium Dichromate Solution
 Tenth normal Volumetric Potassium Permanganate Solution
 Tenth normal Volumetric Potassium Sulphocyanate Solution
 Tenth normal Volumetric Silver Nitrate Solution
 Tenth normal Volumetric Sodium Chloride Solution
 Tenth normal Volumetric Sodium Thiosulphate Solution
 Volumetric Alkali Cupric Tartrate Solution

P G Normal Volumetric Hydrochloric Acid Solution
 Half normal Volumetric Hydrochloric Acid Solution
 Tenth normal Volumetric Hydrochloric Acid Solution
 Hundredth normal Volumetric Hydrochloric Acid Solution
 Normal Volumetric Potassium Hydroxide Solution
 Tenth normal Volumetric Potassium Hydroxide Solution
 Hundredth normal Volumetric Potassium Hydroxide Solution
 Half normal Volumetric Alcoholic Potassium Hydroxide Solution
 Tenth normal Volumetric Iodine Solution
 Tenth normal Volumetric Sodium Thiosulphate Solution
 Tenth normal Volumetric Silver Nitrate Solution
 Tenth normal Volumetric Ammonium Rhodanate Solution

Preparing and Setting Solutions for Volumetric Analysis For the proper performance of Volumetric work it is necessary to employ the following pieces of apparatus —

A burette (preferably Mohr's) fitted with a glass stop-cock, graduated from 0 to 50 c c and subdivided into c c and into $\frac{1}{10}$ c c

Pipettes A series of pipettes graduated to deliver 10, 15, 20, 25, 50 and 100 c c

A graduated glass cylinder, preferably stoppered, graduated at 1000 c c, the intermediate graduations being 100 c c subdivided into 10 c c It should be capable of holding when filled to the zero mark 1000 grammes of Distilled Water at 15 5° C (60° F), and should have preferably an ascending and descending scale

A stoppered glass measuring flask with single graduation, holding when filled to the mark on the neck 1000 c c of Distilled Water, and capable of containing 1000 grammes of Distilled Water at 15 5° C (60° F) This flask is commonly known as a litre flask, though it must be borne in mind that a standard litre represents the volume occupied by 1000 grammes (1 kilogramme) of Distilled Water at 4° C (39 2° F), the temperature of its maximum density, and at a pressure of 760 mm of Mercury 1 millilitre, one thousandth part of this standard litre, is equivalent to 1 00016 c c, or 1 c c is equivalent to 0 99984 millilitre The term 'mille' has been suggested as an abbreviation of a millilitre to more accurately describe the one thousandth part of a litre Unless marked to the contrary, it is assumed that litre flasks, or any similar graduated glass measuring vessels, have reference to the standard litre graduated at the above named temperature [4° C (39 2° F)] The *BP* gives a caution (which will probably be considered unnecessary), to shake the Volumetric Solutions before use to ensure uniformity, and a note regarding their preservation in stoppered bottles, but omits to mention the necessity of ensuring the cleanliness of the bottles and the necessity for washing thoroughly The *USP* directs that all bottles in which Volumetric Solutions are to be kept, as well as the other measuring vessels employed, should before use be thoroughly rinsed in Distilled Water, and then with two or three small portions of the solutions which they are about to contain, and that when not in use the apparatus should be kept

filled with Distilled Water. The *PG* does not give directions for the observance of any special precautions. It is extremely important that great attention should be observed in setting Normal Solutions, as any experimental error is greatly magnified when Fiftieth normal and Hundredth-normal Solutions are prepared from them. These weaker strength solutions should in every case be carefully set before being used for titration, and either adjusted to a strictly correct content, or else a factor ascertained by which the number of c.c. may be converted into those of a strictly correct solution.

VOLUMETRIC SULPHURIC ACID SOLUTION

Normal Solution of Sulphuric Acid should contain in each litre a weight equivalent in grammes to one half its molecular weight, that is to say, it should contain $97.34 - 2 = 48.67$ grammes of pure Hydrogen Sulphate. In preparing the solution a useful method is to carefully weigh out in a clean, dry, stoppered vessel 1 gramme of the Sulphuric Acid to be used in the preparation of the solution, pour it carefully into a small volume of Distilled Water, rinse out the weighing bottle with several small successive quantities of Distilled Water, and determine the weight of pure Hydrogen Sulphate present by titration with Normal Volumetric Sodium Hydroxide Solution, using a few drops of Phenolphthalein as an indicator of neutrality. The Normal Volumetric Sodium Hydroxide Solution is prepared and set in a manner to be hereafter described. Having ascertained the quantity of Hydrogen Sulphate present, the quantity necessary to produce a solution containing a slight excess of the normal amount is weighed out and poured gradually and in small portions at a time into about 500 c.c. of the Distilled Water, the mixture being kept well cooled during the addition. When thoroughly mixed the solution is made up to 1000 c.c., and the mixture well shaken. A portion of the solution is then transferred to a burette, and the exact number of c.c. required to neutralise the molecular equivalent of pure dry Sodium Carbonate determined, Methyl Orange Solution being used as an indicator of neutrality. The balance of the acid remaining in the burette is returned to the original vessel, its volume ascertained, and it is then diluted in the proportion of the number of c.c. which it has actually required to the number of c.c. which it should have required had it been strictly normal. As an example, assuming that the titration of the 1 gramme of Sulphuric Acid has shown that the acid contains 98 p.c. of pure Hydrogen Sulphate, then the quantity necessary to produce a solution containing a little over 48.67 grammes per litre would be the following proportion—as 98 is to 100 so is 48.67 to x , $x = 49.66$ grammes, so that in round numbers about 50 grammes should be weighed out, or, if it be preferred to measure the quantity, about 27 c.c. It should be mixed as described above, and the solution set against the pure dry Sodium Carbonate obtained by igniting pure powdered Sodium Bicarbonate. A quantity of 1.0581 grammes of the pure dry Sodium Carbonate is weighed out, and the Sulphuric Acid Solution added from a burette. The number of c.c. required is noted, and the solution diluted accordingly. Assuming that 950 c.c. of the solution remain after titration, and that x represents the number of c.c. of Sulphuric Acid Solution required to neutralise the above weight of pure dry Sodium Carbonate, it will be x c.c. to be added to the 950 c.c. solution of 950 to 950 + x . In order to ensure accuracy the number of c.c. should be reset.

BP—The *BP* employs a weighed quantity of 50 grammes of Sulphuric Acid, which it dilutes with 900 c.c. of Distilled Water, presumably adding the Distilled Water to the acid, which is a dangerous method of proceeding. It employs titration against pure dry Sodium Carbonate (obtained from the ignition of Sodium Bicarbonate) as a means of setting the acid, and presumably refers to Litmus as an indicator of neutrality, as it inserts a caution to hold off the Carbonic Anhydride. The use of Methyl Orange Solution as an indicator, as described above, obviates the necessity of boiling off Carbonic Anhydride and shortens the time required for the titration.

USP—The *USP* prepares Normal Sulphuric Acid Solution by carefully mixing 80 c.c. of pure concentrated Sulphuric Acid of the *USP* official strength (see p. 80) with sufficient Distilled Water to produce about 1050 c.c., and sets a measured quantity of this strong solution by titration with

Normal Volumetric Potassium Hydroxide Solution, using 2 drops of Methyl Orange Solution as an indicator of neutrality. Having ascertained the number of cc required it proceeds to dilute the strong solution in the proportion of the volume actually required to the proportion which should have been required were the solution of strictly normal strength. As an additional safeguard against error it directs that a second determination of the strength should be made to ensure the exact correspondence of the solution, and if it be still found that solutions differ a new adjustment is directed to be made.

P G—Not included. See Table.

DECI NORMAL VOLUMETRIC SULPHURIC ACID SOLUTION

This solution may be prepared from the Normal Solution by diluting a measured quantity of the stronger solution with sufficient Distilled Water to produce ten times the volume of liquid. Thus 100 cc of Volumetric Sulphuric Acid Solution may be diluted with sufficient Distilled Water to produce 1000 cc of Deci normal Volumetric Sulphuric Acid Solution. It should be carefully set against pure dry Sodium Carbonate in the same manner as described under Volumetric Sulphuric Acid, except that the weight of substance there indicated may be dissolved in Distilled Water and diluted with a further sufficient quantity of Distilled Water to produce 200 cc, 10 cc of this solution may be employed and should correspond exactly to 10 cc of Deci normal Volumetric Sulphuric Acid Solution. In the event of the solutions not strictly corresponding, the number of cc should be noted and the solution diluted accordingly, and should be again reset with a further 10 cc of the pure dry Sodium Carbonate Solution.

B P—The *B P* prepares the Deci normal Solution from the Volumetric Sulphuric Acid Solution in a similar manner to that described above, but does not mention the necessity of setting the finished product.

U S P—The Tenth normal Volumetric Sulphuric Acid of the *U S P* is prepared by diluting 100 cc of Normal Volumetric Sulphuric Acid Solution with sufficient Water to measure 1000 cc at 25° C (77° F). As will be seen from the monographs on the standardised preparations, *e g*, Belladonna, Ipecacuanha, Nux Vomica, etc., this solution is most frequently employed together with Fiftieth normal Volumetric Potassium Hydroxide Solution (*U S P*) in the titration of alkaloids, and in the various instances different indicators of neutrality are used, *e g*, Hæmatoxylin Solution, Cochineal and Iodocsin Solution. The *U S P* authorities have consequently inserted a requirement to the effect that a special experiment should be made, in order to ensure the correspondence of these two solutions towards these indicators of neutrality. In the event of the solutions not corresponding they should be so adjusted that they do exactly correspond.

P G—Not included. See Table.

HALF NORMAL VOLUMETRIC SULPHURIC ACID SOLUTION

Half normal Volumetric Sulphuric Acid Solution may be prepared by diluting a measured volume of Normal Volumetric Sulphuric Acid Solution with sufficient Distilled Water to produce 2 volumes, that is to say 500 cc of the stronger solution may be diluted with sufficient Distilled Water to produce 1000 cc of the weaker solution, and may be set against a solution prepared by dissolving 2.6927 grammes of pure dry Sodium Carbonate in Distilled Water and adding a further sufficient quantity of Distilled Water to produce 100 cc. 10 cc of the Semi normal Volumetric Sulphuric Acid should be equivalent to 10 cc of the Sodium Carbonate Solution, Methyl Orange Solution being employed as an indicator of neutrality. If the solutions do not correspond, the number of cc required should be accurately noted and the corresponding adjustment made, they may then be again re-titrated to ensure strict correspondence.

B P—Not included. See Table.

U S P—The Half normal Volumetric Sulphuric Acid Solution of the *U S P* is prepared on lines similar to those indicated above. The measurements are made at 25° C (77° F). The solution is chiefly used for the titration of the organic salts of Potassium and Sodium, and for this purpose Methyl Orange Solution is usually employed as an indicator of neutrality. The

U S I—A measured quantity of 10 c.c. of Normal Volumetric Potassium should be titrated with the above solution, using 2 drops of Methyl Orange T.S. as an indicator of neutrality, exactly 20 c.c. of the Half-normal Solution should be required. In the event of a divergence between the above solutions, an adjustment should be made in order to ensure their exact correspondence.

P G—Not included. See Table.

FIFTIETH-NORMAL VOLUMETRIC SULPHURIC ACID SOLUTION

This solution may be prepared by diluting 1 volume of Volumetric Sulphuric Acid Solution with sufficient Distilled Water to produce 50 volumes, or 1 volume of Deci-normal Volumetric Sulphuric Acid Solution with sufficient Distilled Water to produce 5 volumes. Thus 20 c.c. of Volumetric Sulphuric Acid Solution may be diluted with sufficient Distilled Water to produce 1000 c.c., or 200 c.c. of Deci-normal Volumetric Acid Solution may be diluted with sufficient Distilled Water to produce 1000 c.c. It may be set against a solution of pure dry Sodium Carbonate prepared by dissolving a weighed quantity of 0.10531 gramme of pure dry Sodium Carbonate in Distilled Water, and adding a further sufficient quantity of Distilled Water to produce 100 c.c. 50 c.c. of this solution exactly corresponds to 50 c.c. of the Fiftieth-normal Solution, Methyl Orange Solution may be used as an indicator of neutrality.

B P—Not included. See Table.

U S P—The *U S P* Fiftieth-normal Volumetric Sulphuric Acid Solution is prepared on similar lines to those above indicated, except that the measurements are made at a temperature of 25° C (77° F). It is chiefly employed for alkaloidal titration, and for this purpose Hæmatoxylin, Cochineal or Iodeosin T.S. are generally employed as indicators of neutrality. The *U S P* does not in this instance require that a special test should be made to ensure correspondence between the solutions with which these indicators are used.

VOLUMETRIC SODIUM HYDROXIDE SOLUTION

A solution of Sodium Hydroxide containing in each 1000 c.c. 39.76 grammes of pure Sodium Hydroxide (NaHO , eq. 39.76). The solutions should be kept in well-corked glass bottles, fitted with a vaselined stopper, an indiarubber stopper, or, preferably, a glass stopper in a tube containing Soda-lime, or it may be kept under a layer of pure liquid.

A convenient method for preparing Volumetric Sodium Hydroxide Solution is to first prepare a concentrated Sodium Hydroxide Solution by dissolving 50 grammes of good stick or good powdered Sodium Hydroxide in 50 c.c. of Distilled Water, the resulting solution is kept under a layer of pure liquid Paraffin. 1 gramme of this concentrated solution is carefully weighed and diluted with about 10 c.c. of Water, a few drops of Phenolphthalein Solution added and the liquid titrated with Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. The number of c.c. of Volumetric Sulphuric Acid Solution required is noted, and the amount of pure Sodium Hydroxide corresponding to this number of c.c. is ascertained by multiplying by 0.03976, having ascertained the percentage of Volumetric Sodium Hydroxide present in each c.c. of the concentrated solution. The solution is removed to represent about 41 or 42 grammes of Sodium Hydroxide, and this quantity is mixed with sufficient Distilled Water to produce 1000 c.c., this will give a solution somewhat greater in strength than the Normal. This solution is set against 20 c.c. of Volumetric Sulphuric Acid Solution, Phenolphthalein Solution being added under that heading, Phenolphthalein Solution being used as an indicator, and the number of c.c. required noted. The balance of the liquid in the burette is returned to the vessel containing the main quantity of liquid. Its volume is carefully ascertained, and is then diluted in the proportion of this number of c.c. to 20 (the number of c.c. which should have been required had the solution been strictly normal), thus, as an example, if it represents the number of c.c. required to neutralise the 20 c.c. of Volumetric Sulphuric Acid Solution and 950 c.c. of liquid are available, it should be diluted in the proportion of x is to 20 so is 950. After this solution has been made the finished product should be checked against Volumetric Sulphuric Acid Solution, and if it be

found that the solution should still not correspond a further adjustment of the solution is necessary until they are in exact agreement. It may be noted that the *B P* employs Litmus and not Phenolphthalein Solution as an indicator of neutrality. It is generally conceded that Phenolphthalein is the more sensitive indicator.

B P—The *B P* Volumetric Sodium Hydroxide Solution is prepared by dissolving 42 grammes of purified Sodium Hydroxide in 1000 c.c. of Distilled Water. It is set by titration against Volumetric Sulphuric Acid Solution, the excessively large quantity of 100 c.c. being employed for the titration, and Litmus is used as an indicator of neutrality. The number of c.c. of the strong solution required to neutralise this quantity is noted, and the solution diluted in the ratio between this number of c.c. and 100. The *B P* makes no requirement that the solution should again be checked after the adjustment has been made, and apparently assumes that the solution is correct.

U S P—Normal Volumetric Sodium Hydroxide Solution (*U S P*) is prepared by dissolving 54 grammes of Sodium Hydroxide (*U S P*) in sufficient Distilled Water to measure 1050 c.c. The solution is standardised by titration against Potassium Bitartrate Solution. The Potassium Bitartrate used is a specially purified Bitartrate obtained from Potassium Bitartrates (*U S P*), the following process being adopted. A weighed quantity of 100 grammes of the salt is digested in a covered beaker on a water bath with a mixture of 85 c.c. of Distilled Water and 25 c.c. of diluted Hydrochloric Acid, the digestion being carried on with intervals of occasional stirring for 3 hours, the mixture is quickly cooled, the solution drained from the precipitate, which is washed by decantation with two successive quantities each of 100 c.c. of Water, the precipitate is collected upon a plain filter, the washing with cold Water is continued until a few drops of the filtrate, when acidified with Nitric Acid, afford no opalescence on the addition of Silver Nitrate T.S. The purified Bitartrate is dissolved in the smallest possible quantity of boiling Water, filtered, and the filtrate constantly stirred whilst being repeatedly cooled. The crystalline precipitate is removed by filtration, when cold washed with 300 c.c. of cold Water and allowed to drain thoroughly. It is then dried until constant in weight at 120° C (248° F). It should be kept in dry, well stoppered glass bottles. A weighed quantity of 9.339 grammes of this salt and 160 c.c. of Distilled Water are introduced into a glass flask, the liquid boiled until solution has taken place and the strong Potassium Hydroxide Solution is added, a few drops (3 to 5) of Phenolphthalein Solution being added as an indicator of neutrality. The number of c.c. of Potassium Hydroxide Solution is noted, and, assuming that this number of c.c. is represented by x , the solution is diluted in the proportion of x to 50. The *U S P* requires that, subsequent to adjustment, the solution should be reset against the solution of a similar quantity of a purified Potassium Bitartrate in a similar manner to that described above, and if still found to be in want of agreement, that a fresh adjustment should be made and the solutions made to strictly correspond. A special precaution is inserted with regard to the liability of solutions of Potassium or Sodium Hydroxide to absorb Carbonic Anhydride from the air, and it is required that these Volumetric Solutions should be preserved in bottles provided with well fitting rubber stoppers, or with tubes filled with Soda lime, which latter provision is also to be observed if the solution is allowed to remain for any considerable time in a burette. The use of Volumetric Sodium Hydroxide in place of Volumetric Potassium Hydroxide is permitted, but as the latter solution is stated to foam less and to attack glass more slowly it has a preference over the former.

P G—Not included. See Table.

DECI-NORMAL VOLUMETRIC SODIUM HYDROXIDE SOLUTION

The Deci-normal Solution may be prepared by diluting 1 volume of Normal Solution with sufficient Distilled Water to produce 10 volumes, that is to say, 100 c.c. of Normal Volumetric Sodium Hydroxide Solution is diluted with sufficient Distilled Water to produce 1000 c.c. The finished solution should be set against a Tenth-normal Volumetric Sulphuric Acid Solution prepared as described under that heading, and the solutions adjusted to correspond, Phenolphthalein Solution may be used as an indicator of neutrality.

The solution should be kept in well-closed glass bottles, preferably fitted with a tube containing Soda-lime, or it may be kept under a layer of pure liquid Paraffin.

BP—The *BP* Deci-normal Volumetric Solution is prepared in a similar manner to the solution described above, but no reference is made to a method or ensuring the accuracy of the finished product, nor is any reference made to precautions to be observed for storage.

$\left. \begin{matrix} USP \\ PG. \end{matrix} \right\}$ Not included See Table

DOUBLE NORMAL VOLUMETRIC SODIUM HYDROXIDE SOLUTION

A solution containing in each 1000 c.c., 79.52 grammes of pure Sodium Hydroxide (NaHO).

$\left. \begin{matrix} BP \\ PG. \end{matrix} \right\}$ Not included See Table

USP—The *USP* prepares this solution by dissolving 90 grammes of Sodium Hydroxide (*USP*) in sufficient Distilled Water to produce about 1000 c.c. The solution is finally set against purified Potassium Bitartrate in a similar manner to that described under Normal Volumetric Sodium Hydroxide Solution (*USP*), and is adjusted so that a measured quantity of 25 c.c. exactly neutralises the quantity of Potassium Bitartrate there mentioned. It is equally important that the same should be observed for keeping this solution, as in the case of the Normal Volumetric Sodium Hydroxide Solution.

NORMAL ALCOHOLIC SODIUM HYDROXIDE SOLUTION

A solution containing 89.76 grammes of pure Sodium Hydroxide (NaHO) in each 1000 c.c. It is prepared on similar lines to the Volumetric Sodium Hydroxide Solution except that Alcohol (90 p.c.) is used in the place of Distilled Water and set against Normal Volumetric Hydrochloric Acid Solution, to be hereafter described, Phenolphthalein Solution being used as an indicator of reaction. It may be adjusted on the same lines as the solution referred to, so that the result may be strictly in agreement, or, preferably, a factor may be calculated, in which the number of c.c. used during a titration may be at once converted into units of strictly Normal Solution.

It is kept in glass bottles provided with well fitting stoppers.

BP—The *BP* information respecting Normal Volumetric Alcoholic Sodium Hydroxide Solution is very meagre. It appears as a footnote to Volumetric Sodium Hydroxide Solution, and officially directs the use of Alcohol (90 p.c.) as a solvent. It may also be presumed that the solution is intended to be standardised in a similar manner to the aqueous solution, but no definite statement to this effect appears. No precautions to be observed for storing the solution are given.

$\left. \begin{matrix} USP \\ PG. \end{matrix} \right\}$ Not included See Table

DECI-NORMAL VOLUMETRIC ALCOHOLIC SODIUM HYDROXIDE SOLUTION

A solution containing 8.976 grammes of pure Sodium Hydroxide (NaHO) in each 1000 c.c. It may be prepared by diluting 1 volume of Normal Volumetric Alcoholic Sodium Hydroxide Solution with sufficient Alcohol (90 p.c.) to produce 10 volumes. It may be standardised against Deci-normal Volumetric Hydrochloric Acid Solution (to be hereafter described) and the solution may be either adjusted to a strictly Deci-normal strength, or a factor may be obtained from which the number of c.c. used during the titration may be converted into strictly Deci-normal strength. This factor is obtained by dividing the number of c.c. which should have been required to react, by the number of c.c. actually required. The solution should be preserved in glass bottles fitted with well-fitting stoppers, or, preferably, with a rubber stopper containing a tube filled with Soda-lime.

BP—The *BP* solution is prepared, presumably, in a similar manner to the Tenth-normal Volumetric Sodium Hydroxide Solution, with the exception that Alcohol (90 p.c.) is used as a diluent in place of Water. No definite directions

are given for the standardisation of the finished solution, nor are any precautions directed to be observed in the storage of the solution. The quantity of solution prepared on each occasion should be only sufficient to last for a short period.

$\left. \begin{array}{l} USP \\ PG \end{array} \right\}$ Not included. See Table

NORMAL VOLUMETRIC POTASSIUM HYDROXIDE SOLUTION

A solution containing 55.71 grammes of pure Potassium Hydroxide (KOH) in each 1000 c.c. It may be prepared by dissolving 100 grammes of good commercial Potassium Hydroxide in sticks in 100 c.c. of Distilled Water. The resulting solution is cooled and kept under a layer of pure liquid Paraffin. A quantity of 1 gramme is weighed out in a weighing bottle, diluted with Distilled Water, and the quantity of Potassium Hydroxide present determined by titration with Volumetric Sulphuric Acid Solution, using a few drops of Phenolphthalein Solution as an indicator of neutrality. Having ascertained the quantity present in 1 gramme, a sufficient quantity is weighed out to yield about 58 grammes, and diluted with sufficient Distilled Water to produce 1000 c.c. The solution is standardised by titration with Volumetric Sulphuric Acid Solution, and adjusted to its requisite strength in a similar manner to the method adopted for Volumetric Sodium Hydroxide Solution. The same precautions are necessary with regard to the storage of the solutions when made.

BP—Not included. See Table.

It is presumed that the *BP* does not intend to include Volumetric Potassium Hydroxide Solution, unless the small type note under alcoholic solutions is intended to refer to both the aqueous and alcoholic solutions, which is scarcely likely. The note accords permission to use an equivalent proportion of Potassium Hydroxide in some instances instead of Sodium Hydroxide.

USP—The *USP* Normal Volumetric Solution is required to contain 55.74 grammes of pure Potassium Hydroxide (KOH) in 1000 c.c. at 25° C (77° F), and is prepared by dissolving 75 grammes of Potassium Hydroxide (*USP*) in sufficient Water to measure about 1050 c.c. It is standardised by methods similar to that described in the *USP* process for setting Normal Volumetric Sodium Hydroxide Solution. The *USP* requires that the strength of the finished solution shall be freshly determined after adjustment has been made, and that if required a readjustment shall be instituted in order that there may be strict correspondence between the solutions. It directs similar precautions to those mentioned under Sodium Hydroxide Solution to be observed in storage.

PG—The Normal Volumetric Potassium Hydroxide Solution of the *PG* is required to contain 56.16 grammes of Potassium Hydroxide in 1 litre. No method is given for its preparation or standardisation, nor are precautions for its storage stated.

DECI-NORMAL VOLUMETRIC POTASSIUM HYDROXIDE SOLUTION

A solution containing 5.571 grammes of Potassium Hydroxide (KOH) in 1000 c.c. It is prepared by diluting 1 volume of Normal Volumetric Potassium Hydroxide Solution with sufficient Distilled Water to produce 10 volumes, *e.g.*, 100 c.c. of Normal Volumetric Potassium Hydroxide Solution is diluted with sufficient Distilled Water to produce 1000 c.c. It may be standardised against Deci-normal Volumetric Sulphuric Acid Solution, and in the event of its being found not to correspond should be readjusted till the solutions are in strict accordance.

It should be kept in glass bottles fitted with well fitting rubber corks, preferably fitted with a Calcium Chloride tube containing fragments of Soda lime, or it may be kept under the surface of a layer of pure liquid Paraffin.

BP—Not included. See Table, see also note under *BP* Volumetric Potassium Hydroxide Solution.

USP—The *USP* solution is prepared on similar lines to that given above, with the exception that the Normal Potassium Hydroxide Solution (*USP*) is required to be freshly standardised, and the measurements are made at 25° C (77° F). The solution is set against purified Potassium Bitartrate. A weighed quantity of 0.9839 gramme should require for neutralisation 50 c.c. of the

Tenth-normal Volumetric Potassium Hydroxide Solution. The *USP* requires the same precaution to be taken in this solution as are recommended under Normal Volumetric Solution.

PG—The *PG* solution is required to contain 5.616 grammes of Potassium Hydroxide in 1 litre, and is prepared by mixing 10 cc of Normal Volumetric Potassium Hydroxide Solution (*PG*) and 90 cc of Water. It is standardised by titration against Decinormal Volumetric Hydrochloric Acid Solution. No precautions are stated for its preservation.

FIFTIETH-NORMAL VOLUMETRIC POTASSIUM HYDROXIDE SOLUTION

A solution containing 1.142 grammes of Potassium Hydroxide (KOH) in 1000 cc prepared by diluting 1 volume of the Normal Solution, or 10 volumes of the Decinormal Solution with sufficient Distilled Water to produce 50 volumes, or 20 cc of the Normal or 200 cc of the Decinormal Solution is diluted with Distilled Water to produce 1000 cc. It should be standardised against the Fiftieth-normal Volumetric Sulphuric Acid Solution described on p. 1296, and in the event of its being found incorrect the solution should be readjusted until the solutions are in strict accord. The same precautions which are observed in dealing with the Normal Solution should be also observed here.

USP—A solution containing 1.148 grammes of Potassium Hydroxide (KOH) in 1000 cc prepared in a similar manner to the above, measurements being made at 25° C (77° F). The *USP* does not in this instance require that the solution should be standardised after preparation in order to ensure its correctness, although it states that it is for use together with Tenth-normal Volumetric Sulphuric Acid Solution in such delicate determinations as the titration of alkaloid residues with Hematoxylin, Cochineal or Iodocyan T.S. as indicators of neutrality. It inserts cautions respecting its preservation which are virtually those given under the Normal Solution, with the addition of a recommendation that the solution should be renewed at frequent intervals.

PG—Not included. See Table.

HUNDREDTH-NORMAL VOLUMETRIC POTASSIUM HYDROXIDE SOLUTION

A solution containing 0.5571 gramme of Potassium Hydroxide (KOH) in 1000 cc, prepared by diluting 1 volume of the Normal Solution or 10 volumes of Decinormal Solution with sufficient Distilled Water to produce 100 volumes. It should be standardised against the Hundredth-normal Volumetric Sulphuric Acid Solution, prepared by diluting the Fiftieth normal Solution with an equal volume of Distilled Water, the solution being in turn standardised against pure Distilled Water.

BP—Not included. See Table.

A Centi-normal Volumetric Sodium Hydroxide Solution. Officially described under *Farmacopoeia Britannica* Liquidum, but is not included in the list of Official Volumetric Solutions. It is stated to contain 0.3976 gramme of Sodium Hydroxide per litre. No method of preparation, standardisation, or preservation is given.

USP—A solution containing 0.5574 gramme of Potassium Hydroxide in 1000 cc, prepared by diluting 10 cc of the Normal or 100 cc of the Tenth-normal Solution with sufficient Water to measure 1000 cc. The measurements are made at 25° C (77° F). The usual notice respecting the precautions to be observed for the preservation of the solution is here omitted, but the solution is recommended to be frequently renewed.

PG—The *PG* solution contains 5.574 grammes of Potassium Hydroxide in 1 litre, and is prepared by mixing 10 cc of the Tenth-normal Solution with 90 cc of Water. It is standardised against Hundredth-normal Volumetric Hydrochloric Acid Solution, to be hereafter described.

NORMAL VOLUMETRIC ALCOHOLIC POTASSIUM HYDROXIDE SOLUTION

Normal Volumetric Alcoholic Potassium Hydroxide Solution is rarely, if ever employed, and is not official in either the *USP* or the *PG*. The *BP* description of it is brief.

It is a solution containing 55.71 grammes of Potassium Hydroxide in 1000 c.c. of Alcohol (90 p.c.). The *BP* permits its use in 'certain' cases instead of Alcoholic Sodium Hydroxide, but does not state what these certain cases are.

SEMI-NORMAL VOLUMETRIC ALCOHOLIC POTASSIUM HYDROXIDE SOLUTION

A solution containing 27.855 grammes of Potassium Hydroxide (KOH) in Alcohol (90 p.c.). It may be conveniently prepared in the following manner. A weighed quantity of good commercial Potassium Hydroxide in sticks is dissolved in about 20 c.c. of Water and titrated with Normal Volumetric Sulphuric Acid Solution, using Phenolphthalein Solution as an indicator. From the results of this titration the percentage of pure Potassium Hydroxide in the sample is calculated, and a sufficient quantity taken to ensure a slight excess over the calculated weight, thus, assuming the sample to have indicated 85 p.c. of the pure Hydroxide, the quantity required for the solution should be as 85 : 100 :: 27.855, corresponding to about 33 grammes. Weigh out about 35 grammes of the specimen, dissolve in 20 c.c. of Water and dilute with sufficient Alcohol (90 p.c.) to produce 1000 c.c. The solution may be standardised against Semi-normal Volumetric Hydrochloric Acid Solution, using Phenolphthalein Solution as an indicator of neutrality. A measured quantity of 50 c.c. of the Semi-normal Volumetric Hydrochloric Acid Solution may be taken and the number of c.c. required to neutralise it carefully noted. The solution may then be diluted with Alcohol (90 p.c.) in the proportion between the number of c.c. actually required, to the number of c.c. which should have been required had the solution been strictly Semi-normal. In the event of the solution being only of little more than Semi-normal strength a good plan is to obtain a factor by which it may be converted into terms of strictly Semi-normal Solution, and to allow the extra strength for deterioration on keeping. The solution will be required to be set freshly at each time of using if employed at more than short intervals.

The same precautions which are observed with the Normal Volumetric Potassium Hydroxide Solution should be observed in the case of this solution, and in addition the glass bottles should be of a dark amber tint.

BP—Not included. See Table.

USP—A solution containing 27.87 grammes of Potassium Hydroxide in 1000 c.c. prepared by dissolving about 40 grammes of Potassium Hydroxide (*USP*) in about 20 c.c. of Water, and adding sufficient Alcohol (94.9 p.c.) to produce 1000 c.c. One day is allowed to elapse for the solution to clear, and the supernatant solution is quickly decanted into another bottle. The solution is standardised against a purified Potassium Bitartrate, using Phenolphthalein T.S. (5 drops) as an indicator of neutrality. A weighed quantity of 1.8678 grammes of the purified Bitartrate representing 20 c.c. of a Semi-normal Solution is employed. The number of c.c. required to exactly neutralise is noted and the solution diluted with Alcohol (94.9 p.c.) in the proportion between this number of c.c. and 20. 20 c.c. of the finished solution being required to exactly neutralise 1.8678 grammes of purified Potassium Bitartrate. An alternative method of standardising the solution is with Half-normal Volumetric Hydrochloric Acid Solution, the dilution being made in a similar manner, the finished solution being required to exactly neutralise an equal volume of the Half-normal Volumetric Hydrochloric Acid, the measurements are made at 25°C (77°F). The *USP* inserts precautions respecting the preservation of the solution, and requires that it should be kept in bottles provided with well fitting rubber stoppers, and that it should be protected from the light. The solution decreases in strength somewhat rapidly, and it is required in addition that blank tests should be performed whenever it is employed in titration.

PG—A solution containing 28.8 grammes of Potassium Hydroxide in 1 litre of Alcohol (90 p.c.). No method of preparation is given. It is standardised against Semi-normal Volumetric Sulphuric Acid Solution. The *PG* describes it as a colourless or but slightly yellow coloured liquid. No precautions respecting its preservation are included.

DECI-NORMAL VOLUMETRIC ALCOHOLIC POTASSIUM HYDROXIDE SOLUTION

A Deci-normal Alcoholic Solution of Potassium Hydroxide is seldom if ever employed in official volumetric work. Neither the *USP* nor the *PG* includes such a solution. The *BP* in the vaguest possible terms as a footnote to Volumetric Sodium Hydroxide Solution. From this description it may be taken to mean a solution containing 5.571 grammes of Potassium Hydroxide in 1000 c.c. of Alcohol (90 p.c.) which is for use in 'certain cases' instead of Sodium Hydroxide Solution.

NORMAL VOLUMETRIC HYDROCHLORIC ACID SOLUTION

A solution containing 36.49 grammes of Hydrogen Chloride (HCl) in 1000 c.c. It may be prepared by a method similar to that described under Volumetric Sulphuric Acid Solution, except that Hydrochloric Acid is substituted for Sulphuric Acid, it may be set against pure dry Sodium Carbonate as there described.

BP—Not included. See Table.

USP—A solution containing 36.46 grammes of absolute Hydrochloric Acid in 1000 c.c. It is prepared by mixing 13.158 at 25° C (77° F)] with sufficient Water to measure 1000 c.c., the measurements being made at 25° C (77° F). It is standardised with Normal Volumetric Potassium Hydroxide Solution, using Methyl Orange T.S. as an indicator of neutrality. The number of c.c. of Normal Volumetric Potassium Hydroxide Solution required to neutralise 10 c.c. of the above solution mixed with about 20 c.c. of Water is noted and the acid solution diluted so that the solutions are in strict accord. After the dilution has been made the *USP* directs that a further trial of its strength should be made, and if still found to be incorrect it should be rejected.

PG—A solution containing 36.46 grammes of absolute Hydrochloric Acid in 1 litre. No directions are given for its preparation or standardisation.

SEMI-NORMAL VOLUMETRIC HYDROCHLORIC ACID SOLUTION

A solution containing 18.095 grammes of Hydrogen Chloride in 1000 c.c. prepared by mixing 1 volume of the Normal Solution with sufficient Distilled Water to produce 2 volumes, e.g., 500 c.c. of the Normal Solution is diluted with sufficient Distilled Water to produce 1000 c.c. It may be standardised by titration against Normal Volumetric Potassium Hydroxide Solution, of which 10 c.c. should exactly neutralise 20 c.c. of the Semi-normal Acid. In the event of its not being so, it should be readjusted and a fresh titration should be made in order to ensure the solutions being strictly in accord.

BP—Not included. See Table.

USP—A solution containing 18.09 grammes of absolute Hydrochloric Acid in 1000 c.c. prepared by diluting 500 c.c. of Normal Volumetric Hydrochloric Acid Solution with sufficient Distilled Water to measure exactly 1000 c.c., the measurements are made at 25° C (77° F). No directions are given to ensure the finished solution being of the correct strength.

PG—A solution containing 18.09 grammes of absolute Hydrochloric Acid in 1 litre. No method of preparation or standardisation given.

DECI-NORMAL VOLUMETRIC HYDROCHLORIC ACID SOLUTION.

A solution containing 3.619 grammes of Hydrogen Chloride in 1000 c.c. prepared by diluting 1 volume of the Normal Solution with sufficient Distilled Water to produce 10 volumes, e.g., 100 c.c. of the Normal Solution is diluted with sufficient Distilled Water to produce 1000 c.c. It may be standardised against Deci-normal Volumetric Sodium Hydroxide Solution, using Phenolphthalein Solution as an indicator of neutrality. In event of the solutions not strictly corresponding, a readjustment should be made in order that they may be in strict accord.

BP—Not included. See Table.

Although not included in the official list of Volumetric Solutions it is used

in the titration of the residual alkaloid from the assay of ~~Liquid Extract~~ of Belladonna and is described under *Extractum Belladonnae Liquidum*, as containing 3.619 grammes of HCl per litre. No method of preparation, standardisation, or preservation is given.

U S P—Not included. See Table.

P G—A solution containing 3.6464 grammes of absolute Hydrochloric Acid in 1 litre prepared by mixing 10 c.c. of the Normal Solution and 90 c.c. of Water. No directions for its standardisation are included.

CENTI-NORMAL VOLUMETRIC HYDROCHLORIC ACID SOLUTION

A solution containing 0.3619 gramme of Hydrogen Chloride in 1000 c.c. prepared by diluting 1 volume of Normal or 10 volumes of Deci-normal Solution with sufficient Distilled Water to produce 100 volumes, *e.g.*, 10 c.c. of Normal or 100 c.c. of Tenth-normal Solution is diluted with sufficient Distilled Water to produce 1000 c.c. The solution is standardised with Centi-normal Volumetric Potassium Hydroxide Solution, and in the event of the solutions not strictly corresponding, a readjustment should be made, so that they may be in strict accord. The solution requires to be carefully standardised as it is used in alkaloidal titrations.

B P }
U S P } Not included. See Table.

P G—A solution containing 0.3646 gramme of absolute Hydrochloric Acid in 1 litre. It is directed to be prepared by mixing 10 c.c. of Tenth-normal Volumetric Hydrochloric Acid Solution and 90 c.c. of Water.

DECI-NORMAL VOLUMETRIC IODINE SOLUTION

A solution containing 12.59 grammes of pure Iodine prepared by the re-sublimation of Iodine answering the official test, mixed with 25 p.c. of its weight of dry Potassium Iodide, the resulting crystals being freed from moisture by drying over fused Calcium Chloride or over Sulphuric Acid in a desiccator. A weighed quantity of 12.7 grammes is mixed with 18 grammes of pure Potassium Iodide (free from Iodate) and dissolved in about 25 c.c. of Water. When completely dissolved the solution is diluted by the cautious addition of Water to measure 1000 c.c. It is standardised against Deci-normal Volumetric Sodium Thiosulphate Solution (to be hereafter described), using, if necessary, Starch Mucilage as an indicator. The solution may also be standardised by titration with pure Arsenious Anhydride. The number of c.c. of the Iodine Solution required to completely react with the Deci-normal Volumetric Sodium Thiosulphate Solution employed is noted, and the solution diluted accordingly (as described under Volumetric Sulphuric Acid Solution), so that it shall be strictly Deci-normal, or a factor may be obtained by dividing the number of c.c. which should have been required, by the number of c.c. actually required, and this factor used to interpret the solution into strictly Deci-normal terms.

It should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from the light.

B P—A solution containing 12.59 grammes of pure Iodine and 18 grammes of pure Potassium Iodide in 1000 c.c. The official method of setting the solution is by titration against Arsenious Anhydride, pure Barium Thiosulphate, or other suitable substance, and it is permitted to either use the solution as it is with a correction for difference in strength between it and Deci-normal, or to convert it into a strictly Deci-normal Solution. The *B P* gives no instructions regarding the preservation of the solution.

U S P—A solution containing 12.59 grammes of pure Iodine in 1000 c.c. of Water. It may be prepared by two methods.—1. By dissolving 12.59 grammes of pure Iodine and 18 grammes of pure Potassium Iodide in 300 c.c. of Water and diluting with sufficient Water to measure exactly 1000 c.c. The measurements are made at 25° C (77° F). The pure Iodine is prepared from Iodine (*U S P*) by careful re-sublimation, first over boiling Water to remove moisture, Cyanogen Iodide, Bromide and Chloride, and then by mixing it with 5 p.c. of its weight of dry Potassium Iodide and re-sublimation on a sand bath, any moisture still adhering is removed by drying over Calcium Chloride. A determination of

the strength of this solution is made at the time of using, unless it is freshly prepared. 2 A weighed quantity of 14 grammes of Iodine (*U S P*) and 18 grammes of Potassium Iodide (*U S P*) are dissolved in about 300 c.c. of Water and the solution diluted to 1000 c.c. The resulting solution is standardised by titration with 10 c.c. of Tenth-normal Volumetric Sodium Thiosulphate Solution, the number of c.c. recorded, the ascertained volume of the solution diluted in the ratio of the number of c.c. actually required to the number of c.c. which should have been required (namely 10). After dilution the strength is again verified to ensure strict accordance between the solutions, and in the event of their still differing a fresh adjustment is made followed by re-determination in order to ensure the solutions being strictly correct.

The solution is directed to be kept in glass-stoppered bottles, but no other precaution specified.

P G—A solution containing 12.685 grammes of Iodine and 20 grammes of Potassium Iodide in 1 litre. The Iodine is directed to be dissolved by the aid of 20 grammes of Potassium Iodide, but otherwise no instructions are given respecting the determination or standardisation of the solution, no directions are included concerning the preservation of the solution when made.

DECI-NORMAL VOLUMETRIC BROMINE SOLUTION

$\left. \begin{matrix} B P \\ P G \end{matrix} \right\}$ Not included. See Table.

The solution is only employed in the determination of Phenol.

U S P—A solution containing 7.936 grammes of pure Bromine in 1000 c.c. It is prepared by dissolving 2 grammes of Potassium Bromate and 50 grammes of Potassium Bromide in sufficient Water to measure 900 c.c. The solution is standardised by means of Potassium Iodide Solution. A measured quantity of 20 c.c. of the above solution is transferred into a 250 c.c. volumetric flask, mixed with 75 c.c. of Water, 5 c.c. of pure Hydrochloric Acid and 5 c.c. of Potassium Iodide T.S. The liberated Iodine is titrated with Decinormal Volumetric Sodium Thiosulphate. The number of c.c. required is noted, and an ascertained volume of the solution diluted in the ratio of the number of c.c. of Tenth-normal Solution actually required to the number of c.c. of Tenth-normal Solution which should have been required. A fresh adjustment of the strength of the solution is made, and in the event of their still differing it is readjusted so that the solutions shall be strictly in accord.

The solution is directed to be kept in well-stoppered bottles of a dark amber tint.

DECI-NORMAL VOLUMETRIC SODIUM THIOSULPHATE SOLUTION

A solution containing 24.644 grammes of crystallised Sodium Thiosulphate ($\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$) in 1000 c.c. of Distilled Water. It may be prepared by dissolving of the crystallised salt in about 250 c.c. of Distilled Water, adding a further quantity of Distilled Water to measure 1000 c.c. It may be set against a weighed quantity of re-sublimed Iodine described under Decinormal Volumetric Iodine Solution, or against a measured quantity of Decinormal Volumetric Potassium Bichromate Solution (as described) and Potassium Iodide Solution. A measured quantity of 20 c.c. of Decinormal Volumetric Potassium Bichromate Solution is introduced into a solution of 1.67 grammes of Potassium Iodide in 50 c.c. of Water, and 10 c.c. of diluted Hydrochloric Acid added. The liberated Iodine is titrated with the above solution, the number of c.c. accurately noted, and an ascertained volume of the approximately Decinormal Thiosulphate Solution diluted in the ratio between the number of c.c. actually required, and the number of c.c. which should have been required. After dilution, the solution is reset in a similar way to that described above. In order to ensure accuracy, and if found not to be in agreement, it should be readjusted so that the solutions should be in strict accord.

It should be kept in well-stoppered glass bottles of a dark amber tint and in a cool atmosphere.

B P—A solution containing 24.644 grammes of crystallised Sodium Thiosulphate ($\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$) in 1000 c.c. of Distilled Water. It is officially directed

to be prepared by dissolving 28 grammes of the salt in 1000 c.c. of Distilled Water, and to be set against Volumetric Iodine Solution, Mucilage of Starch being employed as an indicator. An excessively large quantity of Volumetric Iodine Solution is employed for titration, namely 100 c.c. The solution is diluted in accordance with the result of this determination, and adjusted to contain the above mentioned amount of crystallised Sodium Thiosulphate. No directions are given with regard to the preservation of the solution.

U S I' —A solution containing 24.644 grammes of crystallised Sodium Thio sulphate in 1000 c.c. at 25° C (77° F). It is prepared by dissolving 30 grammes of the salt in sufficient Distilled Water to measure 1000 c.c. The Sodium Thio sulphate employed must be free from Sulphates and Sulphites, free alkalis and Calcium salts. The solution is standardised with Tenth normal Volumetric Potassium Bichromate Solution and Potassium Iodide. The solution is diluted in accordance with the result obtained with this Volumetric determination, and a fresh determination is made to ensure the accuracy of the solution. In the event of the solutions being still found lacking in agreement a fresh adjustment is made in order to render them in strict accord.

The solution is required to be kept in glass stoppered bottles carefully protected from dust.

I' G —A solution containing 24.832 grammes of Sodium Thio-sulphate in 1 litre of Water. No directions are given for its preparation, standardisation, or preservation.

DECI NORMAL VOLUMETRIC SILVER NITRATE SOLUTION

A solution containing 16.869 grammes of Silver Nitrate in 1000 c.c., prepared by dissolving this quantity of the salt in about 250 c.c. of Distilled Water, and diluting the solution with a further quantity of Distilled Water to measure 1000 c.c. It may be standardised with Deci normal Volumetric Hydrochloric Acid Solution or with Deci normal Volumetric Sodium Chloride Solution. A measured quantity of 10 c.c. of either solution may be employed. The number of c.c. required is accurately noted and an ascertained volume of the solution is diluted with Distilled Water in the ratio between the number of c.c. of Deci-normal Volumetric Silver Nitrate Solution actually required, and the number of c.c. which should have been required, namely 10. In order to ensure that the dilution is correct a re determination of the strength of the solution should be made, and in the event of the solutions still not agreeing a further adjustment should be made so that the solutions may be in strict accord.

When finished the solution should be kept in well stoppered glass bottles of a dark amber tint in a cool atmosphere and protected as far as possible from contact with dust and light.

B P —A solution containing 16.869 grammes of Silver Nitrate in 1000 c.c. of Water prepared by dissolving the salt in a sufficient quantity of Water and further diluting with Distilled Water to produce 1000 c.c. The strength of the solution is officially directed to be checked by titration against pure Sodium Chloride or pure Hydrochloric Acid Solution of definite strength, and the solution adjusted according to the result of this Volumetric determination or the real strength of the solution recorded in order that the requisite correction may be made. No directions are contained for the verification of the solution after dilution.

Opaque stoppered bottles are recommended for its preservation, but amber-tinted glass bottles are preferable.

U S P —A solution containing 16.869 grammes of Silver Nitrate in 1000 c.c. of Water prepared by dissolving the pulverised and dried salt, completely dehydrated at 130° C. (266° F.), in sufficient Water to measure 1000 c.c. In this instance it appears to be assumed that the solution will be found correct when made, as no directions are given for its standardisation, although the various methods in which the solution may be employed for titration are recorded. Thus, there is the method of direct titration where the Volumetric Solution is added direct to the solution of a weighed quantity of the salt with Potassium Chromate T.S. as an indicator, there is the Volhard method of indirect titration where an accurately measured excess of the Tenth-normal Volumetric Silver Nitrate Solution is added to the solution to be determined, acidified with Nitric Acid, and the uncombined excess of Tenth normal Volumetric Silver Nitrate Solution is determined

by titration with Tenth-normal Volumetric Potassium Sulphocyanate Solution, using Ferric Ammonium Sulphate T.S. as an indicator, and the method of titration generally employed for the assay of Hydrocyanic Acid and Cyanides where the formation of a permanent red color indicates the end of the reaction.

Preparation of the acid solution in glass-stoppered vials of a dark amber colour, and to be carefully protected from dust and sunlight.

P U—A solution containing 16.977 grammes of Silver Nitrate in 1 litre. No directions as to its preparation, standardisation, or preservation.

VOLUMETRIC POTASSIUM BICHROMATE SOLUTION

A solution containing 4.87 grammes of pure Potassium Bichromate ($K_2Cr_2O_7$), dissolved in Distilled Water, that is to say a solution containing one-tenth of the salt as 1 molecule of Potassium Bichromate liberates 3 atoms of Oxygen which are capable of oxidising 6 atoms of Hydrogen.

It may be prepared by dissolving the above-named quantity of the well-dried and finely powdered salt in about 250 cc of Distilled Water and diluting the solution with a further quantity of Distilled Water, sufficient to make the above volume. It may be standardised against Deci-normal Volume of Sodium Thio-sulphate Solution, using Potassium Iodide Solution and Starch Solution, or by titration with Deci-normal Volumetric Sodium Hydroxide Solution. In the latter case it must be recollected that the solution is Potassium Bichromate. It may also be set against pure Ferrous Ammonium Sulphate Solution. In the latter case a solution containing one-sixth molecular equivalent of Ferrous Ammonium Sulphate corresponds to the Deci-normal solution. If found too strong the solution may be diluted in a similar manner to other volumetric solutions, and re-standardised after dilution.

It should be kept in well-stoppered glass bottles of a dark amber colour and protected as far as possible from contact with dust and direct sunlight.

BP—A solution containing 4.87 grammes of Potassium Bichromate in 1000 cc of Distilled Water. This weight of the salt in about half the required volume of water gives a solution so obtained with a further quantity of Distilled Water to measure exactly 1000 cc. It is officially directed to be standardised by titration with pure Ferrous Ammonium Sulphate Solution, or other reliable substance, and either dilution of the solution to a Deci-normal strength or the ascertainment of its true strength is recommended.

USP—A solution containing 4.8718 grammes of pure Potassium Bichromate in 1000 cc of Distilled Water. It is prepared by dissolving the quantity of pulverised salt dried at $120^\circ C$ ($248^\circ F$) in sufficient measure to exactly 1000 cc at $25^\circ C$ ($77^\circ F$). A method of standardising the solution is not given but references are made to its uses, it is further pointed out that when used with Phenolphthalein as an indicator the solution is Tenth-normal when it contains 14.614 grammes in 1000 cc, but when used as an oxidising agent a solution containing the amount stated at the commencement of the paragraph is the value of a Tenth-normal Solution. When employed for titrating various compounds, it is necessary to use a burette to a solution of the Ferrous Salt in Water. It is necessary with Sulphuric Acid, Potassium Ferri-cyanide T.S. as an indicator, it is also employed in conjunction with Potassium Iodide and Sulphuric Acid in standardising Tenth-normal Sodium Thio-sulphate Solution.

P G—Not included. See Table.

DECI-NORMAL VOLUMETRIC OXALIC ACID SOLUTION

A solution containing 6.255 grammes of pure crystallised Oxalic Acid ($H_2C_2O_4 \cdot 2H_2O$, eq 125.10) in 1000 cc of Distilled Water. It may be prepared by dissolving 6.5 grammes of the crystallised salt in about 100 cc of Distilled Water and adding a further sufficient quantity of Distilled Water to produce 1000 cc. It may be standardised by titration with Deci-normal Volumetric Sodium Hydroxide Solution. A measured quantity of 20 cc of the latter solution is introduced into a porcelain basin, and the Oxalic Acid Solution run in from a burette. Phenolphthalein Solution long used as an indicator of neutrality.

The number of c.c. required is accurately noted, and an ascertained volume of the solution is then diluted in the ratio of the number of c.c. actually required to the number of c.c. which should have been required (20 c.c.) After dilution it is again standardised in order to ensure the correctness of the solution, and if found to be still lacking in agreement the solution should be further adjusted until it is in strict accord. It is employed in standardising Deci normal Volumetric Potassium Permanganate Solution.

B.P. — Not included. See Table.

U.S.P. — A solution containing 6.255 grammes of pure crystallised Oxalic Acid in 1000 c.c. at 25° C (77° F). It may be prepared by dissolving 6.4 grammes of pure Oxalic Acid in sufficient Water to measure 1000 c.c. It is standardised by titration against 10 c.c. of a freshly standardised Tenth normal Volumetric Potassium Hydroxide Solution diluted with twice its volume of Water, Phenolphthalein T.S. being used as an indicator of neutrality, the titration being conducted at a boiling temperature. The solution is diluted in accordance with the usual instructions for the dilution of volumetric solutions. A fresh determination being made to ensure the accuracy of such dilution, and a readjustment being made should the solution be found to be still incorrect, to render them in strict accord. The *U.S.P.* mentions that it deteriorates on standing. It is used for standardising Tenth normal Volumetric Potassium Permanganate Solution.

P.G. — Not included. See Table.

DECI NORMAL VOLUMETRIC POTASSIUM PERMANGANATE SOLUTION

A solution containing 3.1374 grammes of pure crystallised Potassium Permanganate in 1000 c.c. It may be prepared by dissolving 3.5 grammes of the pure crystallised salt in about 250 c.c. of Distilled Water, and when solution is complete, diluting it with sufficient Distilled Water to measure 1000 c.c. It may be standardised by titration with Deci normal Volumetric Oxalic Acid Solution, a measured quantity of 20 c.c. of the latter solution is introduced into a glass flask, mixed with about 1 c.c. of Sulphuric Acid, heated over a Bunsen flame, and the Permanganate Solution run in from a burette until a faint pink colour (permanent for about 30 seconds) is produced. The number of c.c. required is accurately noted, and an ascertained volume of the approximately Deci normal Permanganate Solution is diluted in the ratio between this number of c.c. and 20. To ensure correctness the solution is re standardised with a further quantity of the Volumetric Oxalic Acid Solution, and in the event of the solutions not agreeing a further adjustment is made until the solutions are in strict accord.

The finished solutions should be kept in well stoppered glass bottles of a dark amber colour and protected as far as possible from contact with dust and light.

B.P. — Not included. See Table.

U.S.P. — A solution containing 3.1396 grammes of pure crystallised Potassium Permanganate in 1000 c.c. of Water, measured at 25° C (77° F). It is prepared by dissolving 3.3 grammes of pure crystallised salt by the addition of 1000 c.c. of Distilled Water. The solution being boiled for about 5 minutes, the flask is then plugged with Cotton Wool and the suspended matter allowed to deposit, the clear portion of the solution being poured off, the Water used for dilution is Water that has been distilled over Potassium Permanganate. The solution is standardised against 10 c.c. of an accurately standardised Tenth normal Volumetric Oxalic Acid Solution with the addition of 1 c.c. of pure concentrated Sulphuric Acid, the Permanganate Solution being diluted in the usual manner with sufficient Distilled Water to produce a strictly Tenth-normal Solution. An alternative method of standardisation is also mentioned. A measured quantity of 20 c.c. of the approximate Permanganate Solution is introduced into a solution of about 1 gramme of Potassium Iodide in 10 c.c. of diluted Sulphuric Acid, and the mixture diluted with 200 c.c. of Distilled Water. The liberated Iodine is titrated with an accurately standardised Tenth normal Volumetric Sodium Thiosulphate Solution. The number of c.c. of the latter solution is noted carefully and the solution diluted in the ratio between the number of c.c. of Permanganate Solution which should have been required and the above number. The *U.S.P.* requires that a fresh trial in the manner described immediately

above should be made after dilution, and if necessary a new adjustment should be made to render the between the solutions perfect

P G—Not included

TENTH-NORMAL VOLUMETRIC SODIUM CHLORIDE SOLUTION

This solution is very little used, and is chiefly employed for the titration of Silver salts or for the standardisation of Volumetric Silver Nitrate Solution. The *U.S.P.* describes the method of preparing pure Sodium Chloride, but a very pure salt may be obtained so cheaply that it is questionable whether it is worth while to do so. See Table

U.S.P.—A solution containing 5.806 grammes of pure Sodium Chloride in 1000 c.c. It is obtained by dissolving 5.806 grammes of pure Sodium Chloride in sufficient Water to measure exactly 1000 c.c. at 25° C. (77° F.). No method is indicated for the standardisation of the solution, and when prepared with the pure salt it may be assumed to be correct. See Table

TENTH-NORMAL VOLUMETRIC POTASSIUM SULPHOCYANATE SOLUTION

This solution is also known under the name of Volhard's Solution, and is chiefly used for the indirect titration of the Silver salts, according to Volhard's method. It is mentioned under Tenth-normal Volumetric Silver Nitrate Solution. Ferric Ammonium Sulphate Solution is usually employed as an indicator.

B.P.—Not included. See Table

U.S.P.—A solution containing 9.653 grammes of pure Potassium Sulphocyanate in 1000 c.c. of Water. It is prepared by dissolving 10 grammes of pure crystalline Potassium Sulphocyanate (which is free from Antimony, Arsenic, Cadmium, Copper, Iron, Lead, Zinc) in 1000 c.c. of Water. It is standardised with Tenth-normal Volumetric Silver Nitrate Solution. A measured quantity of 10 c.c. of the latter solution, together with 3 c.c. of Nitric Acid (free from Nitrous Acid), being diluted with about 100 c.c. of Distilled Water. The Volumetric Sulphocyanate Solution is added from the burette until a perceptible reddish-brown tint is acquired. Ferric Ammonium Sulphate Solution is employed as an indicator. The number of c.c. is accurately noted and an ascertained volume of the remaining solution diluted in the ratio between the number of c.c. actually required to the number of c.c. which should have been required. After dilution a fresh determination should be made, using a measured quantity of 50 c.c. of Tenth-normal Volumetric Silver Nitrate Solution, 5 c.c. of Ferric Ammonium T.S., 5 c.c. of Nitric Acid and 200 c.c. of Water, and the solution should be so adjusted that exactly 50 c.c. of the Potassium Sulphocyanate Solution will produce the above-mentioned tint. If necessary a further adjustment may be made until the solutions strictly correspond.

P.G.—Not included. See Table

ALKALINE CUPRIC TARTRATE SOLUTION

Potassium-cupric Tartrate Solution, Alkaline Cupric Tartrate Solution, or Fehling's Solution, consists of two solutions. The first solution contains 69.28 grammes of crystallised Copper Sulphate and 1 c.c. of Sulphuric Acid in 1000 c.c. of Water, and is prepared by dissolving 84.64 grammes of crystallised Copper Sulphate in sufficient Water to effect solution, adding 0.5 c.c. of Sulphuric Acid and sufficient Distilled Water to produce 500 c.c. The second solution contains 50.2 grammes of Sodium Potassium Tartrate and 134 grammes of Sodium Hydroxide in 1000 c.c. of Distilled Water, and is prepared by dissolving 176 grammes of Sodium Potassium Tartrate and 77 grammes of Sodium Hydroxide in a sufficient quantity of Water to effect solution, and diluting with sufficient Distilled Water to produce 500 c.c. When required for use the solution of Copper Sulphate is mixed with an equal quantity of the alkaline Tartrate, the above solution is seldom employed volumetrically. Pavy's Solution, described

on p 468, is more convenient, and is more generally employed in volumetric work

B P—The *B P* solution is on the lines indicated above It is not employed volumetrically in the official volume

U S P—The *U S P* alkaline Cupric Tartrate Solution is employed volumetrically It consists of two solutions —1 The Copper Solution, which is prepared by dissolving 34.67 (more correctly 34.6663) grammes of carefully selected un-effloresced pure Cupric Sulphate, free from adhering moisture, in a sufficiency of Distilled Water, and diluting to measure 500 c c at 25° C (77° F) The solution is directed to be kept in small, well stoppered bottles 2 The Alkaline Tartrate Solution, which is prepared by dissolving 173 grammes of crystallized Potassium Sodium Tartrate together with 75 grammes of Potassium Hydroxide in a sufficiency of Water to measure exactly 500 c c at 25° C (77° F) This solution is directed to be kept in small bottles fitted with rubber stoppers When required for use the solutions are mixed in equal volumes

P G—A weighed quantity of 3.5 grammes of Copper Sulphate is dissolved in 90 c c of Water, and the solution mixed with a solution of 17.5 grammes of Sodium Potassium Tartrate in 30 c c of Water, the latter solution having been previously mixed with 40 grammes of Sodium Hydroxide Solution (15 p c)

TENTH NORMAL VOLUMETRIC AMMONIUM RHODANATE SOLUTION

This solution is employed by the *P G* instead of Tenth normal Volumetric Potassium Sulphocyanate Solution for the determination of the excess of Volumetric Silver Solution in the indirect Silver titration Almost the only instance in which it is used in this volume is in the determination of Volatile Oil of Mustard

B P }
U S P } Not included See Table

P G—A solution containing 7.618 grammes of Ammonium Rhodanate in 1 litre No method of preparation, standardisation, or preservation is indicated

INDICATORS OF NEUTRALITY

The following list shows at a glance the Solutions employed by the three Pharmacopœias as indicators in Volumetric Analysis

<i>B P</i>	Cochineal Tincture	<i>P G</i>	Phenolphthalein T S
	Litmus		Rosolic Acid T S
<i>U S P</i>	Methyl Orange		Starch T S
	Phenolphthalein		Turmeric Tincture
	Starch Mucilage		Hæmatoxylin
	Turmeric Tincture		Iodocsin Solution
	Brazil Wood T S		Litmus paper
	Cochineal T S		Phenolphthalein Solution
	Hæmatoxylin T S		Rosolic Acid Solution
	Iodocsin T S		Starch Solution
	Litmus paper and T S		Turmeric paper
	Methyl Orange T S		Turmeric Tincture

BRAZIL WOOD SOLUTION

Brazil Wood solution is coloured yellow by acids and crimson red by alkalis It is used chiefly as an indicator of neutrality in the titration of alkaloids

B P }
P G } Not included See Table

U S P—A solution obtained by boiling 50 grammes of finely cut Brazil Wood for 30 minutes with 100 c c of Water, the evaporated Water being replaced from time to time The mixture is allowed to cool, strained, 100 c c of the strained liquid are mixed with 25 c c of Alcohol (94.9 p c), the solution filtered It is required that the solution be excluded from contact with ammoniacal vapours,

COCHINEAL SOLUTION.

Cochineal solution assumes a yellow or yellowish-red coloration with acids, and a violet coloration with alkalis. As an indicator it is used chiefly in the titration of alkaloids.

It may be employed as an indicator of neutrality in the titration of solutions containing Ammonia, and affords a useful means of judging the neutrality of Ammonium Acetate and Ammonium Citrate Solutions, *see* Liquor Ammonii Acetatis and Liquor Ammonii Citratis. It may also be used for the titration of inorganic acids.

B P—The solution employed by the *B P* is the official Tincture of Cochineal described on p. 421.

U S P—A filtered solution prepared by macerating 1 gramme of unbroken Cochineal for 4 days with a mixture of 20 c.c. of Alcohol (94.9 c.c.) and 60 c.c. of Water. The *U S P* states that it is useless for titrating organic acids.

P G—Not included. *See* Table.

HÆMATOXYLIN SOLUTION

A solution containing 0.2 p.c. w/v of Hæmatoxylin prepared by dissolving 0.2 of a gramme of Hæmatoxylin in sufficient Alcohol (90 p.c.) to produce 100 c.c. The solution is used as an indicator of neutrality in the titration of alkaloids. It assumes a yellow or orange colour in acid solutions and a violet to purple colour in alkaline solutions. It is advisable in working with this indicator to experiment side by side with an equal quantity of a neutral liquid containing exactly the same amount of the solution as has been added to the liquid under trial or, as very different tints may be assumed according to the volume of the solution used.

B P—Not included. *See* Table.

U S P—A 0.2 p.c. w/v solution of Hæmatoxylin prepared by dissolving 0.2 of a gramme of crystalline Hæmatoxylin in 100 c.c. of Alcohol (94.9 p.c.). The *U S P* specifies about 5 drops are to be used for each titration. The titration is to be considered complete when the change in colour remains permanent upon the addition of 1 drop of the volumetric solution after vigorous shaking.

P G—A solution prepared by dissolving a crystal of Hæmatoxylin in 100 c.c. of Alcohol (90 p.c.). The instructions are to use a solution of this composition as an indicator for each titration.

IODEOSIN

A solution containing 0.1 p.c. w/v of Iodeosin [(Tetraiodfluorescein) $C_{20}H_6I_4O_2$, eq. 829.20] prepared by dissolving 0.1 of a gramme of Tetraiodfluorescein in 100 c.c. of Alcohol (90 p.c.). The solution is colourless in the presence of acids, but assumes a pink coloration in the presence of alkalis. It is an extremely sensitive indicator, and is suitable for the titration of minute quantities of alkaloids. The Alcohol used in its preparation should be absolutely neutral.

B P—Not included.

U S P—A 0.1 p.c. w/v solution of Iodeosin in 100 c.c. of Alcohol (94.9 p.c.) prepared by dissolving 0.1 of a gramme of Iodeosin in 100 c.c. of Alcohol (94.9 p.c.). The *U S P* recommends the volume of the solution titrated should be about 100 c.c. 20 c.c. of Indicator should be added and 5 drops of the Iodeosin T.S., the solution being well shaken after each addition of Volumetric Alkali solution. The titration is continued until a faint pink colour remains after vigorous shaking.

P G—A solution containing 0.2 p.c. w/v of Iodeosin prepared by dissolving 1 part by weight of Iodeosin in 500 parts by weight of Alcohol (90 p.c.). The directions given in the *P G* for the verification of the indicator are as follows. To 100 c.c. of Water contained in a flask of white glass is added sufficient Ether to form a layer 1 cm. in depth, 1 drop of Hundredth-normal Volumetric Hydrochloric Acid Solution and 5 drops of Iodeosin Solution are added, after vigorous shaking the lower aqueous layer should remain uncoloured, but if 2 drops of Hundredth-normal Volumetric Potassium Hydroxide Solution be added, after vigorous shaking the lower aqueous layer should be coloured pale rose.

LITMUS

Litmus is the blue colouring matter prepared from various species of *Roccella*. It occurs commercially in small, dark blue, rectangular cubes, possessing a characteristic floral odour. It is employed in the form of a solution, or as **Litmus paper**. It is unaffected by alkalis, but changes to a red colour on the addition of acid, and this red colour is restored to blue when the solution is again rendered alkaline. It is employed for the titration of acids and alkalis, acid salts, and occasionally in the titration of alkaloids and alkaloidal acid salts, e.g., Quinine Bihydrochloride, but its use in the latter instance is not altogether satisfactory. It yields no satisfactory end reaction with Boric Acid. The titration of Carbonates with this indicator is a tedious process, owing to the necessity of boiling off Carbonic Anhydride before a definite end reaction can be obtained. **Solution of red Litmus** is prepared by the cautious addition of a very dilute Hydrochloric Acid Solution, only just sufficient to ensure a faint red colour being added. Red Litmus paper is prepared by impregnating white bibulous paper with the above solution, and drying.

B.P.—The Litmus of the *B.P.* is obtained from various species of *Roccella*.

U.S.P.—The *U.S.P.* does not mention the source of the Litmus.

P.G.—Not described.

LITMUS SOLUTION

A solution of Litmus may be prepared by repeatedly exhausting the cubes with Water until all soluble matters have been extracted, evaporating the mixed extracts to a small bulk, adding sufficient Acetic Acid to decompose Carbonates, evaporating to a thick extract and adding a large quantity of Alcohol (90 p.c.) which precipitates the blue colour, the latter is washed with hot Alcohol (90 p.c.) and dissolved in Water. **Blue or red Litmus paper** may be prepared by dipping strips of calendered or unsized paper in either the blue or red solution, and drying.

B.P.—A filtered solution of Litmus prepared by exhausting 10 grammes of Litmus for 1 hour with 3 successive portions of Alcohol (90 p.c.), using first 40, 80, and finally 30 c.c. The Litmus (from which all matter soluble in Alcohol of this strength has been removed) is digested in 100 c.c. of Distilled Water.

U.S.P.—A filtered solution of Litmus prepared by exhausting powdered Litmus for 1 hour with 8 separate quantities consisting of about 4 times its weight of boiling Alcohol (94.9 p.c.) in order to remove Elythrotin, the superfluous Alcohol is allowed to drain off and the residue digested with an equal weight of cold Water and filtered. This solution after being acidulated may be used to prepare **red Litmus paper**. The residue is extracted with about 5 times its weight of boiling Water, and after it has been thoroughly cooled, filtered, the filtrate is preserved in wide mouthed bottles closed with a plug of Cotton Wool. The *U.S.P.* states that an addition of 1 drop of Tenth normal Volumetric Acid or Alkali to 50 c.c. of Water containing 5 drops of the indicator should produce a distinct change in colour. The blue and red Litmus paper of the *U.S.P.* is prepared by impregnating strips of white unsized paper, free from wood pulp, with either blue or red Litmus Solution as described above, and drying. Litmus papers should be kept in well stoppered bottles.

P.G.—A solution prepared by digesting 10 parts by weight of Litmus for 24 hours at a temperature of 15° to 20° C (59° to 68° F) with 100 parts by weight of Water, the mixture being repeatedly shaken. The mixture after being allowed to deposit is filtered. It is used for the preparation of **blue Litmus paper**, the following method of procedure being adopted.—Sufficient diluted Sulphuric Acid is added to the above solution, brought to a boiling temperature to cause it to assume only a violet blue coloration when diluted with about 100 parts by weight of Water. Strips of the best unglazed paper are coloured with this 10 p.c. w/w Litmus Solution (neutralised as described above and dried) and are then kept from the light. It is required that blue Litmus paper shall be immediately coloured red by 1 drop of a mixture of 1 c.c. of Tenth normal Volumetric Hydrochloric Acid Solution and 100 c.c. of Water. In the preparation of **red Litmus paper** the above-mentioned Litmus Solution is mixed with sufficient diluted Sulphuric Acid until a test portion diluted with about 100 parts by volume of Water is of a pale red colour, and strips of the best unglazed paper are coloured

with this 10 p c w/w acid Litmus Solution, dried and protected from the light. It is required that red Litmus paper should be immediately coloured blue by the addition of a single drop of a mixture of 1 c c of Tenth-normal Volumetric Potassium Hydroxide Solution and 100 c c of Water. Both blue and red Litmus paper are directed to be preserved in well closed vessels protected from the light.

METHYL ORANGE Pourrier's Orange III

Methyl Orange, Helianthin $\text{NaC}_{11}\text{H}_8\text{N}_2\text{SO}_3$ Pourrier's Tropæolum D

An orange-yellow powder readily soluble in Water, sparingly soluble in Alcohol (90 p c). Commercially it is the Ammonium or Sodium salt of Dimethyl amino-azobenzene-sulphonic Acid or Para-sulphobenzene-azodimethylaniline, a body produced by the action of Dimethylaniline on HNO_3 and H_2SO_4 Acid. The *B P* and the *U S P* employ the Sodium salt. The *U S P* also employs chloric Acid to a hot concentrated aqueous solution. The *U S P* also employs acid, basic Lead Acetate Solution throws down an precipitate, could yield no precipitate with Barium Chloride or Calcium Chloride Solution, nor on the addition of an alkali Hydroxide Solution.

METHYL ORANGE SOLUTION

A solution containing 0.1 p c w/v of Methyl Orange. It may be prepared by dissolving 0.1 of a gramme of Methyl Orange in 50 c c of Distilled Water, adding 10 c c of Alcohol (90 p c), mixing thoroughly and diluting with sufficient Distilled Water to measure 100 c c and filtering. It is used as an indicator of neutrality for the titration of acids and alkalis, and is of special service in the titration of Carbonates, as the latter do not effect the end reaction, and the necessity for boiling off the Carbonic Anhydride is obviated. It is not a satisfactory indicator for use in titration of organic acids, such as Oxalic, Acetic, Citric or Tartaric, as the end reaction is indefinite. It is employed in the titration of alkaloids and alkaloidal salts. Care should be taken that only a minimum quantity of the solution should be used in the titration, the end reaction being more delicate the smaller the quantity of solution employed compatible with the observance of a true end reaction. It affords a yellow coloration with alkalis, and a pink or reddish coloration with acids.

B P—A solution containing 0.2 p c w/v of Methyl Orange in a mixture of Alcohol (90 p c) and Distilled Water, prepared by dissolving 0.2 of a gramme of Methyl Orange in Distilled Water, adding 25 c c of Alcohol (90 p c) and sufficient Distilled Water to make 100 c c.

U S P—A solution 1 p c w/v of Methyl Orange prepared by dissolving 1 gramme of Methyl Orange in sufficient Water to produce 1000 c c. It is mixed with just sufficient Tenth-normal Volumetric Sulphuric Acid Solution to colour the liquid red, and until it just ceases to be transparent and then filtered. The *U S P* directions for carrying out the titrations with this indicator are that 1 to 3 drops are sufficient for the volume of from 50 to 100 c c. It is not to be used in alcoholic or boiling solutions.

P G, *N*, *I*, *C*, *A*

PHENOLPHTHALEIN

Phenolphthalein is described on p 881

PHENOLPHTHALEIN SOLUTION

A solution containing 0.5 p c w/v of Phenolphthalein prepared by dissolving 0.5 of a gramme of Phenolphthalein in 50 c c of Alcohol (90 p c), and diluting with sufficient Distilled Water to produce 100 c c. This affords a convenient strength for use as an indicator of neutrality. It is colourless in acid or neutral solutions, but assumes a fine pink coloration in the presence of even a minute quantity of alkali. It is employed in titrating acids and alkalis. It is the most convenient indicator of neutrality for organic acids, e.g., Acetic, Tartaric, Citric, Oxalic and Valerianic Acids, etc. It is, however, useless for the titration of free Ammonia or Ammonium salts or other fixed alkalis when Ammonium salts are present. It may be used in the titration of Carbonates or

Bicarbonates if the precaution of boiling off Carbonic Anhydride is observed. Its utility in this respect, however, is inferior to Methyl Orange Solution. The great advantage is that it may be used in alcoholic solutions or mixtures of Alcohol and Ether, and many organic acids insoluble in Water may thus be titrated. It may also be used for determining the proportion of acid radical present in alkaloidal salts, the salt may be dissolved in a sufficiency of Distilled Water, sufficient Ether to form a separate layer and to hold the liberated alkaloid in solution is added and titration carried out.

BP—A solution containing 0.2 p.c. w/v of Phenolphthalein prepared by dissolving 0.2 of a gramme of Phenolphthalein in 60 c.c. of Alcohol (90 p.c.), and adding sufficient Distilled Water to measure 100 c.c.

USP—A 1 p.c. w/v solution of Phenolphthalein prepared by dissolving 1 gramme of Phenolphthalein in 50 c.c. of Alcohol (94.9 p.c.) and diluting with sufficient Water to produce 100 c.c. The *USP* employs **Phenolphthalein Paper** which is prepared by soaking white unsized paper in the solution, and drying. The *USP* recommends 3 drops of the solution as a sufficient quantity for use with 50 c.c. of solution to be titrated.

PG—A solution containing 1 p.c. w/w of Phenolphthalein obtained by dissolving 1 part by weight of Phenolphthalein in 99 parts by weight of Alcohol (68 to 69 p.c.).

ROSOLIC ACID SOLUTION

A solution containing 0.2 p.c. w/v of Rosolic Acid, Methylaniline or Corallin, $C_{18}H_{14}O_4$, eq. 301.94.

Rosolic Acid is obtained by the action of Nitrous Acid on Paranitrosaniline or Rosaniline, or by treating a dilute solution of Aniline Hydrochloride with Sodium Nitrite, which results in the formation of Diazoaniline Hydrochloride, which when boiled with the addition of Sulphuric Acid affords Rosolic Acid. The solution is prepared by dissolving 0.2 of a gramme in sufficient Alcohol (60 p.c.) to measure 100 c.c. The colour of the solution, which is pale yellow, is unaffected by acids, but in the presence of alkalis a violet red colour is assumed. It is employed chiefly in the titration of Ammonia and Ammonium salts, and in the titrations involving the use of Sodium or Potassium Sulphites, such as the determination of Citral in Lemon Oil. It is also employed in the determination of Formaldehyde. It is not suitable for the titration of Carbonates, nor, with the exception of Oxalic Acid, for organic acids.

BP—Not included. See Table.

USP—A solution containing 1 p.c. w/v of commercial Rosolic Acid prepared by dissolving 1 gramme of the commercial acid in 10 c.c. of Alcohol (48.9 p.c.), and adding sufficient Water to measure 100 c.c. The quantity recommended by the *USP* for each 100 c.c. of solution to be titrated is 0.5 c.c. The use of Peonin (Aurin, R) is permitted in the place of Rosolic Acid. When used for the titration of ammoniacal solutions the *USP* requires that they should be highly diluted.

PG—A solution containing 1 p.c. w/w of Rosolic Acid prepared by dissolving 1 part by weight of Rosolic Acid in 100 parts by weight of Alcohol (90 p.c.).

STARCH TEST SOLUTION

A solution containing 1 p.c. w/v of Starch prepared by titrating 1 gramme of pure Potato Starch with 10 c.c. of Water, mixing with 30 c.c. of boiling Water, boiling for a few minutes, cooling, diluting with sufficient Distilled Water to produce 50 c.c. and mixing with an equal volume of Glycerin. A solution of this strength, after filtration or decantation from insoluble cell envelopes, will keep bright for years. It is employed as a sensitive reagent for Iodine.

BP—A recently prepared solution containing approximately 2 p.c. of Starch prepared by rubbing 1 gramme of Starch with sufficient Distilled Water to produce a smooth paste, and adding a further sufficient quantity of Distilled Water to produce 50 c.c., boiling for a few minutes, and cooling.

USP—A freshly prepared and filtered solution containing approximately 0.5 p.c. w/v of Starch prepared by titrating 0.5 of a gramme of Starch with

5 c.c. of Water and adding sufficient boiling Water with constant stirring to measure 100 c.c.

P G—A filtered solution of indefinite strength prepared by shaking a small piece of white Starch with hot Water and filtering the solution.

TURMERIC

Turmeric as an indicator is seldom used in the three Pharmacopœias, when required the Tincture is usually employed. Its chief use is for the detection of Boric Acid. Strips of Turmeric paper so immersed in a solution of free Boric Acid in a diluted solution of a Borate that only one half the paper is coloured, yield a brown coloration if Boric Acid is present, changing to a dark green or greenish-black on the addition of a fixed alkali solution, or on the addition of Ammonia Solution.

B P—A Tincture of Turmeric prepared by macerating 1 of bruised rhizome to 6 of Alcohol (90 p.c.). Turmeric paper is unglazed white paper which has been soaked in this Tincture and dried.

U S P—A filtered Tincture prepared by first exhausting a convenient quantity of brown Turmeric root with repeated small quantities of Water, and digesting the dried residue with 6 times its weight of Alcohol (94.9 p.c.) for several days. **Turmeric paper** is white unsized paper impregnated with the Tincture and dried.

P G—A Tincture prepared by digesting at a gentle heat 10 parts of coarsely powdered Turmeric root with 75 parts by weight of Alcohol (90 p.c.) with frequent shaking. After being allowed to deposit, the liquid is filtered. It is used for the preparation of Turmeric paper, and for this purpose 10 parts by weight of the above Tincture are diluted with 30 parts by weight of Alcohol (90 p.c.) and 40 parts by weight of Water and strips of the best unglazed paper are impregnated with this mixture, and whilst protected from the light are dried at the ordinary temperature of the air. **Turmeric paper** should assume a brown coloration on the addition of a single drop of a mixture prepared by mixing 1 part of normal Potassium Permanganate Solution and 2 parts of Water.

It should be kept in well-closed vessels and protected from the light.

THERMOMETRIC MEMORANDA

AND

SPECIAL TESTS

THE *B P* requires that the Thermometers used for the determination of temperatures at which specific gravities, melting points or boiling points are taken should be compared with a standard Thermometer, and any inaccuracy noted, corrections made where necessary and also that the Zero points of the instruments should be verified from time to time. It inserts definite instructions for the determination of the **melting points** of substances mentioned in the Pharmacopœia, but the object of this detailed description is not apparent, as will be seen on page 6 of the Materia Medica, Professor Attfeld having stated that 'in the future it must distinctly be understood that the method described in the *B P Appendix* has not necessarily been the one by which the melting points in the Pharmacopœia have been determined'. In the case of Cera Flava and Ceticum the Pharmacopœia gives specific instructions for the determination of the melting point.

The medium used for raising, to the necessary temperature, the capillary tube in which the melting points are taken is Water if the substance melts below 100° C (212° F), whilst liquid Paraffin, Hard Paraffin or Glycerin are suitable vehicles for substances melting at higher temperatures. The *B P* mentions Sulphuric Acid, but where substances equally suitable are available such a strong mineral acid is undesirable and may be dangerous. The *B P* gives the following formula by which a corrected melting point may be calculated with approximate accuracy from the observed melting point —

$$\text{Corrected temperature} = T + 0.000143 (T - t) N$$

T equals the observed or uncorrected temperature, t equals the mean temperature of the emergent column, N equals the length of the emergent column in scale degrees.

The **boiling point** of the substance is determined in a distillation flask with side leading tube, the bulb of the thermometer is passed through a cork closing the neck of the flask, and, although not actually immersed in the liquid, should be near it, and the whole of the thread of Mercury should, if possible, be surrounded by the vapour. In the case of Amyl Nitrite the *B P* inserts the words, 'the bulb of the thermometer, not dipping below the surface of the residual liquid'. Power (*L B P* '00, 324) states it is not clear why the word residual is inserted when the bulb of the thermometer should not at any time dip below the surface of the liquid.

SPECIAL TESTS

The more important of the special tests which are referred to in the present volume are those adopted for the detection of Arsenic, for which the **modified Gutzert's or Bettendorf's test** is used, the ***U S P* Time-limit test for heavy metals**, the methods for the determination of the **Saponification value** and **Iodine value** of fixed oils, the **Maumené Sulphuric Acid test**, the determination of the **Acid, Ester and Saponification values** of Gums and Gum resins.

THE MODIFIED GUTZEIT'S TEST The *BP* mentions a limit for arsenical contamination only in the monograph on Glycerin depending on Siebold's modification of Gutzeit's test, which is capable of detecting and estimating about 1 part of Arsenic in 250,000 parts of Glycerin. The *USP* describes the test in detail, and the general directions to be followed are contained essentially in the following brief outline: 2 grammes of Zinc, which is required to be free from Arsenic, Sulphur and Phosphorus, which should not contain more than 0.05 p.c. of Iron, and should otherwise correspond to all the *USP* tests for Zinc, 20 c.c. of (8 p.c.) Arsenic-free Hydrochloric Acid and 5 c.c. of Water are introduced into a flask, and first a roll of clean dry gauze and then a piece of test-paper, soaked in a piece of cheese-cloth about 1 decimetre square soaked in the same solution, is introduced into the neck of the flask and pressed with sufficient firmness to retain its place, 1 cm. space should be allowed above the gauze, the top of the flask, after being securely covered by folding over it a piece of pure white filter paper, the centre of which has been moistened with a drop of a saturated alcoholic Mercuric Chloride Solution, and the paper dried, the moistening being repeated with 2 successive drops of the same solution, and in each instance re-dried. The reaction between the Zinc and Hydrochloric Acid is allowed to proceed until the major portion of the Zinc has passed into solution. If no yellow or orange colour appears on the Mercuric Chloride test-paper the materials may be considered sufficiently pure for a trial of a direct test. While the blank test is being carried out, a flask containing a similar mixture of Zinc and 8 p.c. Hydrochloric Acid, together with 5 c.c. of the 1 in 10 solution of the substance to be tested, is prepared, a wad of clean dry gauze, followed by the Lead Acetate test gauze, is introduced, and, after observing the same precautions as above, the Mercuric Chloride test-paper is fastened over the top, after the lapse of at least half an hour, the Mercuric Chloride test-paper is examined for the presence of a yellow stain. A distinct yellow or orange spot indicates the presence of Arsenic, much in excess of 1 in 100,000. In preparing a solution of the chemical for testing, 5 c.c. of the 1 in 10 aqueous solution, or a residue of the residue remaining after special treatment in 5 c.c. of Water, is mixed with 1 c.c. of a mixture of equal volumes of Sulphuric Acid and Water, followed by 10 c.c. of a freshly-prepared saturated solution of Sulphurous Acid, the liquid is heated in a small beaker upon a bath of boiling Water until it is free from excess of Sulphurous Acid, and has been reduced to a volume of 5 c.c. In testing for Phosphorus compounds, *e.g.*, Hypophosphorous Acid, complete oxidation of the sample is necessary before employing the modified Gutzeit's test.

BETTENDORF'S TEST—The *BP* does not mention the Bettendorf test for Arsenic, the general directions contained in the *PG* will be found under the headings of the respective monographs of those chemicals in which it is used as a test. The *USP* adds, 5 c.c. of a saturated absolutely-freshly-prepared Stannous Chloride Solution in pure concentrated Hydrochloric Acid to 5 c.c. of a solution of the prescribed quantity of the substance in pure concentrated Hydrochloric Acid, heats for 15 minutes in a water-bath of boiling Water, allows the tube to stand for one hour. When the tube is examined over a white surface, a brownish tint is observed in the presence of Arsenic beyond the permissible amount. The tube should be viewed from above and compared with a mixture of 5 c.c. each of pure concentrated Hydrochloric Acid and a similarly prepared Stannous Chloride Solution.

THE USP TIME-LIMIT TEST FOR HEAVY METALS—The *USP* has combined a limit of time with the Hydrogen Sulphide test for heavy metals. 10 c.c. of a 1 in 20 aqueous solution of the substance under examination is introduced into a test tube of about 40 c.c. capacity, is acidulated with 1 c.c. of diluted Hydrochloric Acid, warmed to about 50° C (122° F), and an equal volume of freshly-prepared Hydrogen Sulphide T.S. added, the mixture allowed to stand in the well-stoppered test-tube at a temperature of 35° C (95° F) for at least half an hour. At the end of this time any coloration or turbidity is correctly noted, Ammonia Water is added in excess, and the solution again examined for coloration or turbidity, care should be taken to ensure an excess of Hydrogen Sulphide throughout the test. A comparison is made in a test tube containing

under inspection, but without the substance under examination, the observation being made crosswise by a reflected light against a white surface. The 1 in 20 dilution of the substance to be examined has been extended by the Committee of Revision to a dilution of a 1 in 100, for Iron the total dilution is extended to 1 in 300. For chemical substances to be tested for Antimony and Arsenic dilution has not been extended, it remains at 1 in 20. This test detects Antimony, Arsenic, Cadmium, Copper, Iron, Lead and Zinc, if present.

SAPONIFICATION VALUE—The Saponification value represents in milligrammes the weight of pure Potassium Hydroxide required to saponify 1 gramme of the fixed Oil or Fat. The *B.P.* has not yet introduced Saponification values for fixed Oils and Fats. Saponification values are included in both the *U.S.P.* and the *P.G.*, the methods adopted by the latter Pharmacopœia being generally given in the text under the heading of the individual fixed Oil. The general directions for the *U.S.P.* determination are as follows. A weighed quantity of 1.5 to 2 grammes of the purified and filtered Fat is introduced into a flask, together with 25 c.c. of Half normal Volumetric Alcoholic Potassium Hydroxide Solution, a small funnel is placed in the neck of the flask, which is then heated on a water bath for half an hour, 1 c.c. of Phenolphthalein T.S. is added and the excess of the Volumetric Alcoholic Potassium Hydroxide Solution determined with Half normal Volumetric Hydrochloric Acid Solution. A blank test is carried out alongside the determination. The difference between the number of c.c. of Half normal Volumetric Hydrochloric Acid Solution used in the blank test and in the actual determination is multiplied by 27.87 and divided by the weight in grammes of the Fat or Oil.

IODINE VALUE—The Iodine value represents the percentage of Iodine required to combine with the unsaturated fatty acids present in the fixed Oil or Fat. The *B.P.* does not include Iodine values. The general directions followed by the *P.G.* are given under the heading of each individual fixed Oil or Fat. The *U.S.P.* gives the following general directions for the determination of the Iodine value. A weighed quantity of 0.3 of 1 gramme of the fixed Oil or Fat is dissolved in 10 c.c. of Chloroform contained in a bottle of 250 c.c. capacity, a measured quantity of 25 c.c. of a mixture of equal volumes of Alcoholic Iodine T.S. and alcoholic Mercuric Chloride T.S. is introduced, the bottle stoppered, set aside in a cool place protected from the light for 4 hours, an excess of Iodine being ensured throughout this period. A measured quantity of 20 c.c. of Potassium Iodide T.S. is added, and 50 c.c. of Water, the excess of Iodine is titrated with Tenth normal Volumetric Sodium Thio-sulphate Solution, the number of c.c. is noted, a blank experiment made with the same actual quantities of reagents is carried out, the Iodine being determined by the same standard solution. The number of c.c. of Tenth normal Volumetric Sodium Thio-sulphate Solution used in the actual test is deducted from the number of c.c. of Tenth normal Volumetric Sodium Thio-sulphate Solution used in the blank, the difference multiplied by 12.59 and the product divided by 3.

The **alcoholic Iodine T.S.** is prepared by dissolving 25 grammes of Iodine (*U.S.P.*) in 500 c.c. of Alcohol (94.9 p.c.). The **alcoholic Mercuric Chloride T.S.** is prepared by dissolving 30 grammes of Mercuric Chloride HgCl_2 (corresponding to the *U.S.P.* requirements) in 500 c.c. of Alcohol (94.9 p.c.).

Alcohol (90 p.c.) in place of Alcohol (94.9 p.c.) is used in preparing the corresponding solutions of the *P.G.*

MAUMENÉ'S SULPHURIC ACID TEST—Maumene's Sulphuric Acid test indicates the rise of temperature occurring when a weighed quantity of a fixed Oil is mixed with a specified amount of Sulphuric Acid. A weighed quantity of 50 grammes of the oil is introduced into a 7 oz. beaker, which, together with the strong Sulphuric Acid, is immersed in a capacious vessel of Water until they are both of the same temperature, which should not be far from 20° C (68° F). The beaker containing the whole is then wiped and placed in a cotton wool nest, previously made for it in a wider beaker. The thermometer is then immersed and the temperature noted. A measured quantity of 10 c.c. of

concentrated Sulphuric Acid is allowed to run into the oil, the mixture meanwhile being constantly stirred with the thermometer, and about one minute being allowed for the acid to run in, the stirring is continued until no further rise of temperature takes place. The point is readily observed, as the indication remains constant for a minute or two before the temperature begins to fall.

ACID AND SAPONIFICATION VALUES OF GUMS AND GUM-RESINS.—The Acid and Saponification values of Gums and Gum-resins represent the number of milligrammes of pure Potassium Hydroxide required to saponify respectively the free Acid and the free Acids plus the combined Esters of the sample. The Ester value represents the difference between the Acid and Saponification value. The *BP* does not include such figures. Where such figures are given in the *USP* the method of their determination generally appears under each individual subject. The figures to determinations made in the author's laboratory refer to the method given by Dieterich in his *Analysis of Resins, Balsams and Gum resins*.

SPAS IN BRITAIN

- BATH** (Somersetshire) —The only true thermal Water in England. Saline, 21 grains in 20 oz. Chiefly Calcium Sulphate and small quantities of Sodium Sulphate and Magnesium Chloride, with Carbonic Acid gas and Nitrogen. Several baths varying in temperature from 88° to 120° F (31.1° to 48.8° C). For chronic rheumatism, gout, and paralysis. The Water is aerated and sold in bottles under the name of **Sulis Water**. Radium has been discovered in the Waters of Bath and Buxton.
- BUXTON** (Derbyshire) —A Water containing only 2½ grains of salts in 20 oz., with Carbonic Acid gas and Nitrogen. Temp. 82° F (27.7° C). For chronic gout and rheumatism.
- DROITWICH** (Worcestershire) —Mineral. Chiefly Sodium Chloride, about 27½ grains in 20 oz., along with Sodium and Calcium Sulphates. Warm baths are given, usually at a temperature of from 95° to 101° F (35.6° to 38.3° C). Used in muscular rheumatism, sciatica, and in chronic rheumatic and gouty affections.
- FLITWICK** (Amphill, Beds) —Chalybeate, aperient, 31 grains in 20 oz. Iron Carbonate, Magnesium and Sodium Sulphates, Magnesium Chloride and Calcium Carbonate.
- HARROGATE** (Yorkshire) —Several springs, sulphur and chalybeate. The old sulphur spring contains 137 grains in 20 oz., chiefly Chlorides, with Sulphuretted and Unsulphuretted Hydrogen. Of the chalybeate Waters, the new spring contains 62 grains in 20 oz. chiefly Chlorides with about 1½ grains Iron Chloride, together with Carbonic Acid gas and Nitrogen.
- LEAMINGTON** (Warwickshire) —‘Old well,’ saline about 10½ grains in 20 oz., chiefly Sodium and Calcium Chlorides, with Sodium Sulphate and Carbonic Acid. The saline chalybeate Waters contain about 13½ grains in 20 oz., chiefly Calcium, Magnesium and Sodium Chlorides, with Sodium Sulphate and a small quantity of Iron Carbonate. In stomach and liver complaints, in gouty and rheumatic affections.
- LLANDRINDOD WELLS** (Wales) —Mineral, mineral sulphur and weak chalybeate waters. The first contains Sodium Chloride (about 30 to 40 grains in 20 oz.) along with Calcium and Magnesium Chlorides. The second, in addition to being weakly mineralized, contains from 1 to 14 volumes per 1000 of Hydrogen Sulphide gas, and the third about the same amount of Sodium and Calcium Chlorides as the first, along with a small amount of Iron Carbonate. Used in tonic dyspepsia, constipation, and in chronic rheumatism and rheumatoid arthritis.
- LLANGAMMARCH** (Wales) —Barium Water. About 38 grains in 20 oz., chiefly Sodium, Calcium and Magnesium Chlorides, with about ½ grain of Barium Chloride. Cardiac tonic. In glandular affections, gout and rheumatism.
- MALVERN** (Worcestershire) —A table Water nearly pure, containing about ½ grain of mineral salts in 20 oz. Useful in kidney and bladder affections.
- STRATHPEFFER** (Ross-shire) —Two springs, Upper and Lower Sulphurous. Containing chiefly Sodium and Calcium Sulphates, with Sulphuretted Hydrogen. The Upper about 18 grains in 20 oz., the Lower about 18½ grains and slightly less Sulphuretted Hydrogen than the Upper.
- WOODHALL** (Lincolnshire) —About 190 grains in 20 oz., chiefly Sodium, Calcium and Magnesium Chlorides, with Sodium and Potassium Bromides and Potassium Iodide. A ‘mutterlange’ is also used. Useful in chronic rheumatism, scrofula, syphilis, etc.

SPAS — FOREIGN

ACHSELMANNSTEIN (Bavaria) — Saline, aperient, chalybeate. About 2237 grains in 20 oz., chiefly Sodium and Magnesium Chlorides, with Sodium and Calcium Sulphates and Carbonic Acid gas. Baths and Vapor Baths for incipient tuberculosis, cutaneous diseases and uterine disorder. May to September.

ADELHEIDSQUELLE (Heilbrunn, Bavaria) — Saline, about 58 grains in 20 oz., chiefly Sodium Chloride (44 grains), with Sodium Iodide $\frac{1}{4}$ grain and Bromide about $\frac{1}{2}$ grain and Carbonic Acid gas. Acts on the glandular, lymphatic, and cutaneous systems. May to September. Imported.

AESCLAP (Buda-Pesth, Hungary) — Aperient, antacid. About 334 grains in 20 oz. Chiefly Sodium Sulphate (125 grains) and Magnesium Sulphate (154 grains), with Sodium Chloride and Calcium Sulphate. For habitual constipation and disorders of the liver. Imported.

AIX-LA-CHAPELLE (Rhine Province, Germany) — Several springs. Thermal 113° to 133° F (45° to 56.1° C). Saline, sulphurous. About 39 grains in 20 oz., chiefly Sodium Chloride (25 grains) and Sodium Carbonate (about 5 grains), with varying quantities of Sodium Sulphide. Used in cutaneous diseases, rheumatism and syphilis. Summer season April to October. Winter season November to April. Imported.

AIX-LES-BAINS (Savoy, France) — Two chief springs are 'Sulphur Spring' and 'Alum Spring' to 112° F (42.7° to 44.4° C). About 3 $\frac{1}{2}$ grains in 20 oz., chiefly Sodium and Calcium Sulphates, with Sulphuretted Hydrogen in the 'Sulphur Spring'. Rheumatism, gout, eczema. It contains the organic substance 'Glairine' or Bârgine peculiar to sulphur Waters. April to November.

ALET (Aude, France) — Alkaline. Thermal 82° F (27.7° C) for baths, and a feruginous Water 50° F (10° C). Weak in minerals, about 4 $\frac{1}{2}$ grains in 20 oz. Tonic in debility and dyspepsia.

ALEXANDERBAD (Bavaria) — Chalybeate. About 3 grains in 20 oz., of which about $\frac{1}{2}$ grain is Iron Carbonate, with Carbonic Acid gas. May to October.

ALEXISBAD (Germany) — Chalybeate. 'Alexis-Brunnen' and 'Freundschafts-Brunnen' are used for drinking, and the 'Solke-Brunnen' for bathing, June to September.

ALLEVARD (Isere, France) — Gaseous. Iodo-sulphuretted. About 195 grains in 20 oz. Chiefly Calcium and Magnesium Carbonates, Chlorides and Sulphates, with about 0.05 grain of Iodine, Carbonic Acid gas and Sulphuretted Hydrogen. June to September.

APENTA (Hungary) — Aperient. Chiefly Sodium Sulphate (about 161 grains) and Magnesium Sulphate (about 182 grains), with Sodium Chloride and Calcium Sulphate in 20 oz. Imported.

APOLLINARIS (Neuenahr, Rhine Province, Germany) — Alkaline, gaseous. About 22 grains in 20 oz., chiefly Sodium Carbonate (about 11 grains), Chloride and Sulphate, with Magnesium Carbonate. Free Carbonic Acid gas. Imported and drunk as a table Water.

ARABELLA (Germany) — Aperient. Contains chiefly Magnesium and Sodium Sulphate, Sodium Sulphate and Magnesium Chloride. For liver and kidney complaints, gastric catarrh, diabetes.

ARNSTADT (Germany) — Brine spring, for baths. About 2200 grains in 20 oz., of which 2150 grains are Sodium Chloride. Used for scrofula. 'Riedquelle' with about 34 grains Sodium Chloride in 20 oz., for drinking. April to September.

AUTEUIL (France) — Chalybeate. About 28 grains in 20 oz., about 6 grains being Iron and Aluminium Sulphates,

BADEN (Vienna) Saline and sulphurous. About 17 grains in 20 oz, chiefly Calcium, Potassium and Sodium Sulphates, with Magnesium Sulphide and Chloride. Principally used for bathing, also mineralised mud cataplasms in rheumatism. May to October.

BADEN-BADEN (Germany) —Several thermal springs, 124° to 150° F (51° to 65 5° C) 'Hauptquelle' most used for drinking. Saline, about 27 grains in 20 oz. The Lithia Waters contain about 3 grains Lithium Chloride in 20 oz. Other salts chiefly Chlorides and Carbonates of Sodium, Calcium and Magnesium, with trace of Iron Arsenate. For rheumatoid arthritis, chronic gouty affections and paralysis. May to October.

BAGNÈRES-DE-LUCHON (France) —Thermal sulphur springs, 61° to 152° F (16° to 66 6° C). About 2 grains of Iron, Manganese and Sodium Sulphides in 20 oz. Used in chronic cutaneous eruptions, catarrhal diseases of the respiratory organs, etc. June to October.

BARÈGES (Hautes Pyrénées, France) Thermal sulphurous. Temp 81° to 111° F (27 2° to 43 8° C). About 2 grains in 20 oz, chiefly Sodium Sulphide Sulphate and Chloride, with Baryte similar to Haulmo. 'Tun boui' spring used internally, about ½ grain Sodium Sulphide per 20 oz. Skin diseases and chronic rheumatism. June to September. Imported.

BATTAGLIA (Province of Venice, Italy) —Thermal. Four springs. Temp 136° to 160° F (57 7° to 71 1° C). The Waters contain about 13 grains of Sodium Chloride in 20 oz, and are similar to, but more weakly mineralised than those of Baden Baden. Mud baths are also employed, the mud 'Fango' is exported. They are used in chronic gout and rheumatism, and in rheumatoid arthritis. May to October.

BELLTHAL (Rhine Province, Germany) —Alkaline, table Water. About 11 grains in 20 oz, chiefly Sodium, Potassium, Magnesium and Calcium carbonates. Imported.

BERKA (Weimar, Germany) —Chalybeate and sulphurous springs. About 27½ and 16½ grains of solids respectively in 20 oz, chiefly Calcium Sulphate and Carbonate, with about 0 4 grain Iron Carbonate in the chalybeate Water. Used for chronic rheumatism, anæmia and debility.

BETHESDA (Wisconsin, U S A) —About 5 3 grains in 20 oz, chiefly Calcium and Magnesium Carbonates. Used in treatment of kidney diseases.

BILIN (Bohemia) —Gaseous, alkaline. About 47 grains in 20 oz, chiefly Sodium Carbonate (about 29 grains) and Sulphate with Calcium and Magnesium Carbonates. Taken for indigestion and acidity. Also drunk as a table Water. May to September. Imported.

BIRMENSTORF (Switzerland) —Aperient. Temp 50° F (10° C). About 279 grains in 20 oz, chiefly Magnesium Sulphate (about 191 grains) and Sodium Sulphate (about 61 grains), with Calcium Sulphate and other salts. Imported.

BIRRESBORN (Rhine Province, Germany) —Alkaline, gaseous spring. About 43 grains in 20 oz, chiefly Sodium Bicarbonate (about 24 grains), with Sodium Sulphate and Magnesium Bicarbonate. Free Carbonic Acid gas. Table Water.

BOCKLET (near Kissingen) —Chalybeate. Temp 50° F (10° C). Three springs, varying in mineral strength, contain Sodium Chloride and Sulphate, with Calcium and Magnesium Carbonates and about 0 8 grain Iron Carbonate in 20 oz, and much free Carbonic Acid gas. Tonic, useful for anæmic and debilitated patients. May to September.

BONIFACIUS (Hesse Nassau, Germany) —About 122 grains in 20 oz, chiefly Sodium Chloride (about 89 grains), Magnesium Chloride, Calcium Sulphate, Lithium Chloride (about 2 grains), and Magnesium Bromide and Iodide. Stimulates the intestines and urinary organs.

- BONNES** (Basses Pyrénées, France) (**Eaux Bonnes**)—Thermal Temp 72° to 90.5° F (22.2° to 32.2° C) Saline and sulphurous From about 5 to 6 grains in 20 oz, chiefly Sodium and Potassium Chlorides, with Calcium Sulphate, Sodium Silicate and 'Barégine,' with Sulphuretted Hydrogen Used in chronic bronchitis, pharyngitis, and catarrhal affections of respiratory organs June to September
- BORCETTE** or **BURTSCHIED** (near Aix-la Chapelle)—Springs similar to those of Aix-la-Chapelle
- BOURBOULE, LA** (Puy-de-Dôme, France)—Two chief springs, 'Perruie' and 'Choussy' Aisenical, equal to about 0.16 gram of Sodium Arsenate in 20 oz, also Sodium Chloride and Bicarbonate, about 24 grains of each Used in affections of the respiratory organs May to September Imported
- BRIDES-LES-BAINS** (Savoy, France)—Mineral sulphated springs About 16 grains of Sodium Chloride, 10 grains of Sodium Sulphate, with Calcium and Magnesium Sulphates, and minute quantities of Iron and Arsenic Tonic with laxative action in large doses June to September Imported (both Salts and Water)
- BRUCKENAL**, Γ - - ybcate About 0.09 gram Iron Carbonate in 20 oz
- BRUCOURT** (Calvados, France)—Chalybeate About 0.43 gram Iron Carbonate and 4.3 grains of Magnesium Sulphate in 20 oz along with Calcium Bicarbonate Used in anemia and chlorosis
- BUDA-PESTH**—Several springs of bitter Water, such as Hunyadi-János, Apenta, and Franz Josef, *q v* Buda or Ofen (opposite Pesth, Hungary) Thermal Temp 141.5° F (61° C) Internally and for bathing, chiefly Sodium Sulphate and Carbonate In gastric catarrh, obstinate rheumatism and eczema
- BUFFALO LITHIA** (Mecklenburg Co, Va, U S A)—Three springs Most important is No 2, which contains about 12 grains in 20 oz, chiefly Calcium Sulphate and Bicarbonate, Potassium Carbonate (about $3\frac{1}{2}$ grains), with Lithium Bicarbonate (about $2\frac{1}{2}$ grains), Sulphuretted Hydrogen and Carbonic Acid gas
- BUSSANG** (Vosges, France)—Alkaline, ferruginous, mild laxative About 13 grains in 20 oz, chiefly Sodium, Calcium and Magnesium Carbonates, with about 0.08 to 0.1 gram Iron Carbonate Imported
- CAMBRUNNEN** (Taunus)—Gaseous, antacid table Water Imported
- CAPVERN** (Hautes-Pyrénées, France)—Thermal Temp 70° to 76° F (21.1° to 24.4° C), chiefly Calcium Sulphate, about 9 grains in 20 oz Used in gout and gravel
- CARABANA** (Spain)—Aperient Chiefly Sodium Sulphate, about 875 grains in 20 oz Imported
- CARLSBAD** (Bohemia)—Various springs Thermal Hottest is 'Sprudel' Temp 162.5° F (72.2° C) About 21 grains Sodium Sulphate, about 10 grains Sodium Bicarbonate, and about 9 grains Sodium Chloride in 20 oz, with Calcium Carbonate and Carbonic Acid gas For constipation, liver affections, gout, rheumatism, diabetes April to September Imported (both Salts and Water)
- CAUTERETS** (Hautes-Pyrénées) - - Thermal From 103° to 123° F (39.4° to 53.8° C) About 20 oz, of which about 0.1 gram is Sodium Sulphide May to October Imported
- CHALLES** (Savoy, France)—Cold sulphur Waters About 11 grains in 20 oz, of which about 4 grains are said to be Sodium Sulphide In chronic bronchitis, catarrh of throat, and scrofula June to October Imported
- CHARLOTTENBRUNNEN** (Silesia)—Chalybeate. About 7 grains in 20 oz, chiefly Sodium, Calcium, and Iron Carbonates

- CHÂTELDON** (France) — A gaseous, alkaline table Water Imported
- CHÂTEL-GUYON** (Puy de Dome, France) — Alkaline 'Source Gublet' About 72 grains in 20 oz, chiefly Calcium Bicarbonate (18 grains) Sodium Bicarbonate, Magnesium and Sodium Chlorides, with Iron Bicarbonate and Carbonic Acid gas May to October
- CONDAL** (Rubiñat, Pyrenées, Spain) — Aperient About 450 grains in 20 oz, chiefly Sodium Sulphate (about 390 grains), with Magnesium Sulphate (27 grains), Sodium Chloride and Calcium Sulphate Useful in chronic indigestion and affections of the liver and spleen Imported
- CONDILLAC** (France) — A gaseous, alkaline drinking Water Imported
- CONTREXÉVILLE** (Vosges, France) — Several springs, principal is 'Source Pavillon' About 13 grains Calcium Sulphate 3 grains Calcium Bicarbonate in 20 oz, with minute quantities of Iron, Arsenic and Calcium Fluoride For urinary affections and chronic cystitis May to October Imported
- DAX** (Landes, France) — Thermal Temp 88° to 140° F (31° to 60° C) Used for baths and douches in chronic rheumatic affections 'Mud baths' are also given for rheumatism, scurvy and nervous affections
- DESAIGNES** (France) — Alkaline From 27 to 36 grains Sodium Bicarbonate in 20 oz
- DRIBURG** (Westphalia, Germany) — Chalybeate About 50 grains in 20 oz, chiefly Calcium Bicarbonate (12 grains) and Sulphate (9 grains), with about $\frac{1}{2}$ grain Iron Bicarbonate, and much free Carbonic Acid gas May to October
- EILSEN** (Germany) Sulphurous 'Julianenbrunnen' contains about 97 grains in 20 oz, chiefly Calcium Sodium and Magnesium Sulphates, with Sulphurated Hydrogen about 2.5 c m In gout rheumatism and paralysis May to September
- EMS** (Germany) — Alkaline, mineral, thermal Temp from 80° to 120° F (26.6° to 48.8° C) Several springs 'Kirchenbrunnen,' 'Kesselbrunnen,' 'Fürstenbrunnen,' 'Neuequelle,' all contain about 33 grains in 20 oz, chiefly Sodium Bicarbonate (18½ grains) and Chloride (about 9 grains), with Calcium and Magnesium Bicarbonates and over 500 vols Carbonic Acid gas per 1000 In diseases of mucous membranes, catarrh of larynx and bronchi, gout, dyspepsia, cystitis Imported (Water, Salts and Pastilles)
- ENGHIEN** (Paris) — Sulphurous, containing both Calcium Sulphide and Sulphurated Hydrogen For drinking and bathing Imported
- EVIAN-LES-BAINS** (Savoy, France), — Alkaline, table Waters About 2½ grains in 20 oz, chiefly Calcium Carbonate
- FACHINGEN** (Hesse Nassau, Germany) — Alkaline About 47½ grains in 20 oz, chiefly Sodium Bicarbonate (about 35 grains) and Calcium and Magnesium Bicarbonates, with Sodium Chloride The spring is rich in Carbonic Acid gas For acidity in the stomach, and in kidney and bladder diseases Used also as a table Water Imported
- FIUGGI** (Italy) Alkaline About 0.6 grain each of Magnesium Carbonate and Potassium Nitrate in 20 oz, along with Calcium Carbonate, Sodium Chloride, Oxygen, Carbonic Anhydride and Nitrogen Used in gastric catarrh, liver complaints and stomacal affections
- FRANZENSBAD** (Bohemia) — Several springs, varying considerably in mineral constituents 'Franzensquelle,' 'Salzquelle,' 'Wiesenquelle' and 'Kalter Sprudel' are for drinking, and contain Sodium Sulphate (24 to 31 grains in 20 oz), with Sodium Carbonate, and Chloride and Iron Carbonate in varying quantities The Chalybeate 'Moor baths' are baths containing peat Used in rheumatism and gout May to September
- FRANZ JOSEF** (Buda Pesth) — Aperient About 216 grains each of Sodium and Magnesium Sulphates in 20 oz, with Magnesium Chloride, Calcium Sulphate and Sodium Chloride Imported

- FRIEDRICHSHALL** (Saxe-Meiningen) —Aperient According to Liebig, contained about 237 grains in 20 oz chiefly Sodium Chloride (76 Sulphate (58 grains), Magnesium Chloride and Calcium Sulphate of the stomach, liver and urinary organs Imported.
- GASTEIN** (Austria) —Several thermal springs Temp from 78.5° to 121° F (26° to 49.4° C) About 2½ grains in 20 oz, of which almost 2 grains are Sodium Sulphate Chiefly used for bathing For nervous affections May to September
- GEILNAU** (Hesse-Nassau) —Alkaline, mineral Table water
- GEROLSTEIN** (Rhine Province, Germany) —Table Water About 19 grains in 20 oz, chiefly Sodium, Calcium and Magnesium Carbonates
- GISSHUBLER** (near Carlsbad in Bohemia) —Table Water About 20 grains in 20 oz, chiefly Sodium, Calcium and Magnesium Bicarbonates
- GODESBERG** (Rhine Province, Germany) —Chalybeate, gaseous 'Old' spring contains about ½ grain Iron Bicarbonate in 20 oz 'New' spring only used for bathing, about 0.4 grain
- GRIESBACH** (Baden) —Chalybeate, gaseous About 0.6 grain Iron Bicarbonate in 20 oz, with Sodium Sulphate and Calcium Bicarbonate
- GUBER** (Siebenik, Bosnia) —Ferruginous and arsenical About 6½ grains in 20 oz, chiefly Ferric Sulphate (about 3 grains), with Aluminum Sulphate, and about 0.05 grain of Arsenious Acid
- HALL** (Upper Austria) —Mineral Water of 'Tassilloquelle' It contains Sodium Chloride, about 105 grains in 20 oz, along with about 0.002 p.c. of Magnesium Bromide and Iodide respectively The bath salt is used in scrofulous affections The Water is exported and Iodwasser May to September
- HOMBURG** (Hesse-Nassau, Germany) —Laxative, slightly tonic For drinking, 'Elizabeth-Brunnen,' 'Kaiser-Brunnen,' 'Ludwig-Brunnen,' 'Luisen-Brunnen,' and 'Stahl-Brunnen' Varying mineral constituents, all are rich in Carbonic Acid gas Chiefly Calcium Carbonate, Iron Carbonate and Chloride, with Iron Carbonate Useful in gouty affections, dyspepsia, hysteria, anæmia and debility May to September Water and Salts imported
- HUNYADI-JANOS** (Buda-Pesth) —Aperient Chiefly Sodium Sulphate (about 153 grains in 20 oz), Magnesium Sulphate (about 150 grains), with Sodium Chloride Habitual constipation Imported
- ISCHIA** (Italy) —Principal spring, 'Gugitello' Thermal Temp 131° to 149° F (55° to 65° C) About 52 grains in 20 oz, chiefly Sodium Chloride and Bicarbonate, with Carbonic Acid gas Useful in rheumatism, paralysis, skin diseases, etc Spring and Summer
- ISCHL** (Austria) —Bine baths May to September
- JODBAD LIPIK** (Slavonia, Hungary) —Alkaline Thermal 117° F (63.8° C) About 28 grains in 20 oz, chiefly Sodium Bicarbonate, with about ½ grain of Sodium Iodide and Carbonic Acid gas For catarrhal affections of mucous membranes, gout, rheumatism, etc
- JOHANNIS** (Hesse-Nassau) —Alkaline, table Water Containing chiefly Sodium, Calcium and Magnesium Bicarbonates, with Sodium Chloride
- KISSINGEN** (Bavaria) —Principal spring, 'Rakoczy' A mineral water in 20 oz, chiefly Sodium Chloride (about 54 grains), with Magnesium Chlorides, Calcium Carbonate and Magnesium Sulphate Also 'Pandur-Quelle,' similar, and 'Max-brunnen' weaker gas Kissingen bitter Water from 'Soole' springs For constipation, liver affections, catarrhal conditions of stomach and bowels May to September Imported (Salts and Water)

- KOSEN** (Saxony) —About 477 grains in 20 oz, chiefly Sodium Chloride (420 grains) Baths, in scrofula
- KÖNIGSDORFF-JASTRZEMB** (Silesia) —Saline About 109 grains in 20 oz, chiefly Sodium Chloride, with a little Magnesium Bromide and Iodide
- KRANKENHEIL** (Bavaria) —About 7 grains in 20 oz, chiefly Sodium Bicarbonate and Chloride In scrofulous skin diseases May to October
- KREUZNACH** (Rhine Province, Germany) —Several springs 'Elizabethquelle,' chiefly used for drinking About 117 grains in 20 oz, chiefly Sodium Chloride (about 90 grains), with Calcium and Magnesium Chlorides and a little Bromide and Iodide The mother lye, from which the common salt has been crystallised, about 3100 grains in 20 oz, a large amount being Calcium Chloride Tonic to lymphatic system In syphilis, skin diseases, rheumatism and paralysis Water, Salt and Brine all imported
- KRONDORF** (Bohemia) —Table Water About 21 grains in 20 oz, chiefly Sodium, Calcium and Magnesium Bicarbonates, with Carbonic Acid gas In gout and as a diuretic Imported
- KRONENQUELLE** (Oberhalbbrunn, Silesia) —Alkaline About 20 grains in 20 oz, chiefly Sodium Bicarbonate (about $7\frac{1}{2}$ grains), with Calcium and Magnesium Bicarbonates, Sodium Sulphate, Lithium Carbonate and Carbonic Acid gas In nephritic and arthritic affections, and in gouty diathesis May to September Imported
- KRONTHAL** (Germany) —Table Water Chiefly Sodium Chloride, with Calcium Carbonate and Carbonic Acid gas Imported
- LABASSÈRE** (Hautes Pyrénées) —Sulphurous Chiefly Sodium Chloride, with about $\frac{1}{2}$ grain Sodium Sulphide in 20 oz Bronchial and laryngeal catarrh June to October
- LANDECK** (Silesia) —Thermal 66° to 94° F (18° 8' to 20° C) Under 2 grains in 20 oz, chiefly Sodium Bicarbonate and sulphate, with traces of Sulphide and Sulphuretted Hydrogen For bronchial catarrh Also 'Moor baths,' for rheumatism
- LANGENBRUCKEN** (Baden) —Chiefly Sodium, Magnesium and Calcium Sulphates, with Carbonic Acid gas and traces of Sulphuretted Hydrogen For hæmorrhoidal conditions, bronchial irritation and rheumatism
- LEUK or LOÈCHE-LES-BAINS** (Switzerland) —Thermal 102° to 124° F (48° 8' to 51° 1' C) About 18 grains in 20 oz, chiefly Calcium Sulphate, with Magnesium Sulphate In chronic skin affections June to September
- LEVICO** (Austrian Tyrol) —Arsenical and Ferruginous Two strengths, 'strong' and 'mild' The strong contains about 0.07 grain Arsenious Anhydride, with about 33 grains Iron salts in 20 oz, the mild about 0.008 grain Arsenious Anhydride, with about 8 grains of Iron salts June to September Imported
- LIPPIK** —See Jodbad Lipik
- LIPPSPRINGE** (Westphalia, Germany) —'Ammunquelle' contains about 21 grains in 20 oz, chiefly Calcium and Sodium Sulphates In bronchial irritation and tuberculosis May to September
- LUCCA** (Italy) —Thermal 98° to 129° F (36° 6' to 53° 8' C) About 11½ grains in 20 oz, chiefly Calcium and Magnesium Sulphates, with Sodium Chloride Baths in gout and rheumatism June to September
- LUHATSCHOWITZ** (Moravia, Austria) —Several springs Vincenz, Amand and Johann Brunnen are the chief Contain in 20 oz from about 27 to 39 grains of Sodium Carbonate and 21 to 39 grains of Sodium Chloride, with Calcium Carbonate, also Sodium Iodide and Bromide, and Carbonic Acid gas In bronchial, gastric and uterine catarrh, congested liver and hæmorrhoids May to September
- MARCOLS** (France) —Alkaline 21 to 23 grains Sodium Bicarbonate in 20 oz

- MARIENBAD** (Bohemia) — Chief springs are 'Kruza brunnen' and 'Ferdinand-brunnen'. The first about 92 grains in 20 oz. Second about 102 grains in 20 oz., chiefly Sodium Sulphate (41 to 45 grains), Sodium Bicarbonate (14 to 16 grains), with Sodium Chloride, and Calcium and Magnesium Carbonates. Laxative. Useful in obesity, dyspepsia, and chronic constipation. May to September. Imported (Salts and Water).
- MEINBERG** (Germany) — Several springs, varying in strength. Contain Sodium and Magnesium Sulphates, with Calcium Sulphate and Carbonate. Sulphurous mud-baths are used. For scrofula, rheumatism and gout, facial neuralgia, and generally tonic. May to September.
- MERGENTHEIM** (Wurtemberg) — Aperient. About 119 grains Sodium Chloride, 33 grains Sodium Sulphate, and about 22 grains Magnesium Sulphate in 20 oz., with Carbonic Acid gas. For chronic constipation, catarrh of stomach and intestines, etc.
- MONDORF** (Luxembourg) — Mineral, for drinking and bathing. Temp 77° F (25° C), chiefly Sodium Chloride about 75 grains in 20 oz., with Calcium Chloride and Sulphate, and Magnesium Bromide and Chloride.
- MONT DORÉ** (France) — Source 'Madelme' and source 'Bardon,' mostly used internally. About 18 grains in 20 oz. Thermal Waters, temp up to 113° F (45° C), used for baths, drinking, inhalations, etc. For chronic laryngitis and bronchitis. June to September.
- NAUHEIM** (Germany) — 'Kur brunnen' and 'Kais brunnen,' chiefly for drinking. Containing Sodium Chloride 87 to 130 grains in 20 oz. and Calcium Chloride, with Carbonic Acid gas. The bath Waters are about double this strength. Temp 82° to 95° F (27° to 35° C). In cardiac affections, etc. A special form of treatment is adopted here, known as the 'Nauheim method.' May to September.
- NENNDORF** (Germany) — Sulphurous. 'Trinkquelle' only one used for drinking. About 45 grains in 20 oz., chiefly Sodium Chloride and Calcium Sulphate, with Calcium Chloride and Magnesium Chloride. 'Rodenbarg' brine considerably stronger and used for bathing. For rheumatism, gout, cutaneous affections and catarrh of respiratory organs. May to September.
- NEUENAUH** — See **APOLLINARIS**.
- OBERSALZBRUNN** (Silesia) — Alkaline. Chief spring 'Oberbrunnen,' containing about 19 grains Sodium Bicarbonate in 20 oz., with Sodium Sulphate, Magnesium and Calcium Bicarbonates and a small quantity of Lithium Bicarbonate. In nephritic affections and catarrh of bladder. May to September. Imported.
- OREZZA** (Corsica) — Gaseous, chalybeate. A kind of ferruginous, Seltzer Water agreeable to drink. Chiefly Calcium Carbonate, with about 1 grain of Iron Carbonate in 20 oz.
- PFÄFFERS** — See **RAGATZ-PFÄFFERS**.
- PLOMBIÈRES** (Vosges, France) — Thermal. 77° to 155° F (25° to 68° C). About 24 grains in 20 oz., chiefly Sodium, Calcium and Magnesium Sulphates, with Sodium Sulphate. Principally used as baths. In treatment of gastric, dyspepsia, and catarrhal enteritis.
- POLAND** (U.S.A.) — Alkaline. Contains Calcium Carbonate (1.228 grains in each U.S. gallon) along with Magnesium and Sodium Carbonates, Sodium Chloride and Potassium Sulphate. Used in chronic dyspepsia and liver congestion.
- POUGUES** (Loire, France) — Alkaline (calcareous). 'St. Jeger' spring contains about 15 grains Calcium Bicarbonate and 6 grains Sodium Bicarbonate in 20 oz., with Magnesium Bicarbonate and Chloride. Used in dyspepsia, chronic diarrhoea, and urinary affections.
- PULLNA** (Bohemia) — Saline, purgative. About 310 grains in 20 oz., chiefly Sodium and Magnesium Sulphates, with Magnesium Chloride and Carbonate, and Carbonic Acid gas. In obstinate constipation. Imported.

PYRMONT (Waldeck) - Several springs, chalybeate and mineralized 'Hauptquelle' and 'Helenen quelle' are the two chief chalybeate springs used for drinking. They contain from $\frac{1}{2}$ to $\frac{3}{4}$ grain Iron Carbonate in 20 oz. The mineralized Waters contain varying quantities of Sodium Chloride from 63 grains in 'Trinkquelle' to 288 grains in 'Bohllochquelle' in 20 oz. In anæmia, debility, scrofula and functional nervous affections.

RAGATZ-PFAFFERS (Switzerland) Thermal Temp 98° F (36.6° C) About 3 grains in 20 oz, chiefly Magnesium and Sodium Carbonates, with Sodium and Calcium Sulphates. Rich in Nitrogen. June to September.

RAKOCZI See KISSINGEN

RECOARO (Venetia) - Chalybeate. About 25 grains in 20 oz, chiefly Calcium Sulphate and Carbonate, Magnesium Sulphate and about $\frac{1}{2}$ grain Lithium Carbonate, with Carbonic Acid gas. May to October.

REICHENHALL (Bavarian Alps) - Numerous saline springs, most important being 'Edelquelle,' which contains about 2237 grains in 20 oz, of which about 2150 are Sodium Chloride (chiefly used for baths in scrofula, catarrh of the respiratory organs, etc. May to September).

RÉNAISON (France) - Table Water containing Sodium and Calcium Bicarbonates.

RHENS (Rhine Province, Germany) - Mineralized, alkaline, table Water.

RIPPOLDSAU (Prussia) - Three springs used for drinking, 'Josephs quelle,' 'Leopolds quelle,' 'Wenzelquelle.' Chalybeate. About 33 grains in 20 oz, chiefly Calcium Bicarbonate, Sodium Sulphate and Magnesium Sulphate, with about $\frac{1}{2}$ grain Iron Bicarbonate. For anæmia, also useful in pulmonary catarrh, 'Natrone' and 'Schwefelnatrone' artificially carbonated and sulphated to counteract tendency to constipation. May to September.

ROISDORF (Rhine Province, Germany) - A mineralized, alkaline table Water. Chiefly Sodium Chloride and Sodium Carbonate.

ROSBACH (near Homburg, Germany) - A mineralized, alkaline table Water. Chiefly Sodium Chloride and Calcium Carbonate. Imported.

ROYAT (Puy de Dôme, France) - Alkaline Thermal Temp 68° to 95° F (20° to 35° C). Several springs, 'Eugène,' 'César,' 'St Mark,' 'St Victor.' Source 'Eugène' most highly mineralized, contains about 48 grains in 20 oz, chiefly Sodium Bicarbonate and Chloride, with Calcium Bicarbonate, and Carbonic Acid gas. For gout, uric acid diathesis, dyspepsia, chronic laryngitis and bronchitis. May to September. Imported.

RUBINAT (Pyrenees, Spain) - Natural purgative Water. About 909 grains in 20 oz, chiefly Sodium Sulphate (about 840 grains) and Magnesium Sulphate (about 28 grains), with Sodium Chloride and Calcium Sulphate. For constipation, congestion, gastric fever, etc. Imported.

SAINT BOËS (Basses Pyrenees, France) - Sulphurous, bituminous. About 14 grains in 20 oz, Sodium Sulphide with Sulphuretted Hydrogen, Iodine and Arsenic. For bronchitis, laryngitis, and in pulmonary tuberculosis.

SAINT GALMIER (France) - Alkaline table Water, containing Sodium, Calcium and Magnesium Bicarbonates, and may be obtained charged with additional Carbonic Acid gas. Imported.

SAINT GERVAIS (France) - Thermal Temp 102° to 108° F (38.8° to 42.2° C). Three springs, 'Source de Mey,' 'de Gontard,' 'du Torient.' Contain Sodium Sulphate, Sodium Chloride and Calcium Sulphate. In cutaneous affections, chronic rheumatism and dyspepsia. June to September.

SALIES-DE-BÉARN (Basses Pyrenees, France) - Bine baths, containing about 1925 grains Sodium Chloride, with Magnesium and Potassium Chloride, in 20 oz.

SALINS-LES-BAINS (Jura, France) - Saline. About 198 grains Sodium Chloride in 20 oz, with about $\frac{1}{2}$ grain Potassium Bromide, and traces of Sodium Iodide. For scrofulous affections. May to September.

- SALVATOR** (Fpenn., Hunnry) Alkaline, effervescent. With about $\frac{1}{2}$ grain Sodium Bicarbonate, 8 grains Magnesium Bicarbonate and 15 grains Calcium Bicarbonate in 20 oz. For urinary affections, etc.
- SANKT-MORITZ** or **SAINT MORITZ** (Upper Engadine, Switzerland) - Three springs, 'Alt', 'Neu', and 'Piusellusquelle' and the recently discovered 'Sper', all are chiefly chalybeate. About 23 grains in 20 oz, chiefly Calcium, Magnesium and Sodium Bicarbonates, with Sodium Sulphate and about $\frac{1}{4}$ grain Iron Carbonate. June to September.
- SARATOGA** (U.S.A.) - Alterative. Chiefly Sodium Chloride, Calcium Bicarbonate, Magnesium Bicarbonate, with Sodium Iodide and Bromide. Useful in glandular and visceral obstructions and in skin diseases. 'Congress' and 'A' springs are bottled for export.
- SAUERBRUNNEN** (Goshim Hutz, Germany) - A natural mineral, table Water, containing chiefly Magnesium Bicarbonate and Sulphate.
- SCHINZNACH** (Switzerland) - Strongly sulphurous. Thermal. 82.4 to 95° F (28° to 35° C). Chiefly Sodium Sulphate with Potassium and Magnesium Chlorides, and about $\frac{1}{4}$ grain Calcium Sulphide in 20 oz. Rich in Sulphurated Hydrogen. For chronic eczema and all skin eruptions, gout and rheumatism. May to September.
- SCHLANGENBAD** (Nassau, Germany) - Simple thermal Water. Temp 81.5° to 89° F (27.2 to 31.6° C). About $\frac{1}{2}$ grains in 20 oz, chiefly Sodium Chloride. Rich in Oxygen and Nitrogen. Useful in nervous irritability, is stated to have a sedative and beneficial influence on the skin. June, July and August.
- SCHWALBACH** (Nassau, Germany). Several springs. 'Stahlbrunnen' and 'Weimbrunnen' mostly used internally. 'Palmbrunnen' and 'Rosenbrunnen' are used for baths. Chalybeate. Chiefly Magnesium and Calcium Bicarbonates, with $\frac{1}{2}$ to $\frac{1}{4}$ grain Iron Bicarbonate in 20 oz. and excess of Carbonic Acid gas. For anemia and leucorrhoea. June, July and August.
- SEIDLITZ** (Bohemia) - Bitter aperient. About 140 grains in 20 oz, chiefly Magnesium Sulphate (about 100 grains) with Sodium Sulphate and Calcium Sulphate and Carbonate.
- SELTERS** (N) - A mineralised effervescent table Water containing chiefly Sodium Bicarbonates and excess of Carbonic Acid gas. Imported.
- SODEN** (Nassau, Germany) - Several springs. 'Milch-', 'Warm', 'Wilhelms' and 'Sool-brunnen'. Saline, containing chiefly Sodium Chloride from 22 to 140 grains in 20 oz, with Calcium and Magnesium Carbonates, and from 0.2 to 0.7 grain Iron Carbonate. For chronic laryngitis, bronchitis, gout and scrofula. May to September.
- SPA** (Belgium) - Several springs, principal are 'Pierrefontaine' and 'Prince de Condé'. Chalybeate, alkaline. About 40 grains in 20 oz, chiefly Magnesium, Calcium and Sodium Carbonates, with 0.6 to 1 grain Iron Bicarbonate. In anemia, menorrhagia and debility. May to October.
- TARASP** (Switzerland) - Several springs. 'Lucius' and 'Ementia' are sulphated, 'Bonifacius' is chalybeate. They contain about 15 grains Sodium Sulphate, 36 grains Sodium Bicarbonate, 32 grains Sodium Chloride in 20 oz, with other salts. 'Bonifacius' contains about 0.4 grain Iron Carbonate in 20 oz. Carbonic Acid gas. Useful in obesity, gout, rheumatism and skin diseases, anaemia, etc. June to September.
- TAUNUS** (Frankfort) - A muriated, alkaline, table Water, containing chiefly Sodium Chloride, with Calcium Carbonate and excess of Carbonic Acid gas.
- THONON** (France) - Weakly mineralised Waters, similar to those of Evian les-Bains.

TCHITLI (Turkey) —About 55 grains in 20 oz, chiefly Sodium Bicarbonate

TOEPLITZ or **TEPLITZ** (Bohemia) —Alkaline, thermal Temp 83° to 111° F (28.3° to 45° C) —About 6 grains total solids in 20 oz, chiefly Sodium Bicarbonate about 3½ grains —Used in rheumatism, gout, paralysis and nervous affections —May to September

VALS (France) —Several springs varying in mineral strength from 27 grains to 77 grains in 20 oz —Saint Jean, 'Précieuse,' 'Désirée,' 'Rigolette,' and 'Madelme' are the sources mostly used in England —The Waters contain chiefly Sodium Bicarbonate, with Calcium and Magnesium Bicarbonates, and Carbonic Acid gas —In dyspepsia and gastric catarrh —May to October —Imported

VERNET (Pyrenées Orientales, France) —Thermal Sulphur springs, 90° to 154° F (32.2° to 67.7° C) —About ½ grain Sodium Sulphide in 20 oz —Used for drinking and bathing —For rheumatism, cutaneous eruptions and affections of respiratory organs

VICHY (France) —Alkaline, thermal Temp from 57° to 106° F (13.8° to 41.1° C) —Numerous springs, 'Glande Grille,' 'L'Hopital,' 'Celsus,' 'Hauterive,' 'Saint Yorre' —They contain from 70 to 80 grains in 20 oz, chiefly Sodium Bicarbonate from 35 to 45 grains, with Sodium Chloride and other salts —Used in kidney diseases, diabetes, gouty, hepatic and urinary diseases —For drinking and bathing —May to October —Imported (Pastilles, Salts and Water)

VILLACABRAS (Spain) —Aperient Water, contains chiefly Sodium Sulphate

VITTEL (Vosges, France) —Calcareous springs resembling those of Contrexeville —May to September —Imported

WEILBACH (Nassau, Germany) —Two springs 'Schwefelquelle' and 'Nation lithionquellc' —The first, a sulphur Water, contains about 14 grains total solids in 20 oz, with Sulphuretted Hydrogen —The other about 25 grains in 20 oz, chiefly Sodium Chloride and Bicarbonate, with a small quantity of Lithium Bicarbonate —For hæmorrhoids, gout, rheumatism and urinary complaints —Imported

WIESBADEN (Nassau, Germany) —Several springs, the principal being 'Kochbrunnen' —Saline Thermal 100° to 156° F (37.7° to 69° C) —About 79 grains in 20 oz, chiefly Sodium Chloride (about 65 grains), with Calcium and Magnesium Chlorides —In chronic gout and rheumatism, catarrh of larynx and bronchitis —'Wiesbadener Gichtwasser,' a preparation made from Kochbrunnen with the addition of about 70 grains Sodium Bicarbonate in 20 oz —Imported

WILDEAD (Wurtemberg) —Thermal Temp 91.5° to 104.5° F (33° to 40° C) —Numerous springs —About 4 grains in 20 oz, chiefly Sodium Chloride —Baths are used in chronic rheumatism and gout and paraplegic paralysis of lower extremities —June to September

WILDUNGEN (Waldeck, Germany) —Several springs —Principal are 'Helenenquelle' and 'Georg Victor quelle,' containing varying quantities of minerals, chiefly Calcium, Magnesium and Sodium Bicarbonates —Used in cystitis, pyelitis, renal and vesical disorders

WITTEKIND (Halle, Saxony) —Saline Water for drinking, containing about 8½ p c salt —Also mixed with mother lye for baths

CLASSIFICATION OF MINERAL WATERS

Comparatively Free from Salts

Buxton
Evian-les-Bains
Gastein
Malvern
Schlangenbad
Thonon
Wildbad

Saline

Achselmannstern
Adelheidsquelle
Aix-la-Chapelle
Arnstadt
Baden
Baden-Baden
Bath
Battaglia
Bonifacius
Bonnes
Boicette, or Buttscheid
Brides-les-Bains
Buda, or Ofen
Carlsbad
Droitwich
Hall
Homburg
Ischia
Ischl
Kissingen
Kosen
Koenigsdorf-Jastzemb
Krankenheil
Kreuznach
Leamington
Llanganmaich
Mergentheim
Mondorf
Nauheim
Pyrmont
Reichenhall
Salies-de-Béarn
Salins-les-Bains
Saratoga
Selters
Soden
Spa
Wiesbaden
Wittekind.
Woodhall.

Bitter Saline

Aesculap.
Apenta
Arabella
Burmenstorf
Buda-Pesth,
Carabana
Condal
Franz Josef,

Friedrichshall
Hunyadi
Marienbad
Mergentheim
Pullna
Rubinat
Seidlitz
Villacabras

Alkaline, and Gaseous Alkaline

Bellthal
Bilin
Briesboin
Bussang
Chatel-Guyon
Ems
Fischingen
Franzensbad
Ischia
Jodbad Lipik
Johannis
Kiondorf
Kronenquelle
Luhatschowitz
Marcols
Marienbad
Obersalzbrunn
Reisdorf
Rosbach
Royat
Saint Galmier
Salvator
Selters
Tchitli
Toeplitz
Vals
Vichy
Wildungen

Alkaline and Calcareous

Bothesda
Buffalo Lithia
Capvern
Contrezeville
Fruggi
Lappspinge
Lucca
Poland
Pougues
Vittel

Arsenical

Bouiboule
Brides-les-Bains
Bussang
Guber
Levico
Mont Doré.
Vals.

Chalybeate

Alet
 Alexanderbad
 Alexisbad
 Auteuil
 Berka
 Bocklet
 Brides les Bains
 Bruckena
 Buecourt
 Bussing
 Charlottenbrunnen
 Driburg
 Flitwick
 Franzensbad
 Godesberg
 Griesbach
 Guiber
 Harrogate
 Islandrindod
 Olczna
 Pyrmont
 Recoaro
 Rippoldsan
 Saint Moritz
 Schwalbach
 Soden
 Spa
 Tarasp

Sulphurous

Aix-la Chapelle
 Aix les Bains
 Allevard
 Baden
 Bagnères de Luchon
 Barèges
 Beika
 Bonnes
 Cantorets
 Challes
 Eilsen
 Enghien
 Harrogate
 Labassere
 Islandrindod
 Landeck
 Meinberg
 Nenndorf
 Saint Boes
 Stathpeffor
 Schminnach
 Vernet
 Weilbach

Lithiated

Baden Baden
 Bonifacius
 Buffalo Lithia
 Franzensbad
 Kissingen
 Kronenquelle

Obersalzbrunn
 Weilbach

Thermal Springs

Aix la Chapelle, 113° to 133° F (45° to 56° C)
 Baden Baden, 124° to 150° F (51 1° to 65 5° C)
 Bagnères de Luchon, 61° to 152° F (16 1° to 66 6° C)
 Barèges, 81° to 111° F (27 2° to 45 8° C)
 Bath, 88° to 120° F (31 1° to 48 8° C)
 Bittigha, 136° to 160° F (57 7° to 71 1° C)
 Bonnes, 72° to 90 5° F (22 2° to 32 2° C)
 Buda, or Ofen, 141 5° F (61° C)
 Buxton, 82° F (27 7° C)
 Capvern, 70° to 76° F (21 1° to 24 1° C)
 Carlsbad, 'Sprudel,' 162° F (72 2° C)
 Cantorets, 103° to 126° F (39 4° to 53 3° C)
 Dax, 88° to 140° F (31° to 60° C)
 Ems, 80° to 120° F (26 6° to 48 8° C)
 Gastein, 78° to 121° F (26° to 49 4° C)
 Ischia, 131° to 149° F (55° to 65° C)
 Jodbad Lipik, 147° F (63 8° C)
 Landeck, 66° to 84° F (18 8° to 29° C)
 Leuk, or Loeche les Bains, 102° to 124° F (48 8° to 51 1° C)
 Lucca, 96° to 129° F (36 6° to 53 8° C)
 Mondoif, 77° F (25° C)
 Mont Dore, up to 113° F (45° C)
 Nanheim, 82° to 95 5° F (27 7° to 35° C)
 Plombières, 77° to 155° F (25° to 68 3° C)
 Ragatz Pfaffen, 98° F (36 6° C)
 Royat, 68° to 95° F (20° to 35° C)
 Saint Gervais, 102° to 108° F (38 8° to 42 2° C)
 Schminnach, 82 4° to 95° F (28° to 35° C)
 Schlangenbad, 81 5° to 89° F (27 2° to 31 6° C)
 Toeplitz, 83° to 114° F (28 3° to 45 5° C)
 Vernet, 90° to 154° F (32 2° to 67 7° C)
 Vichy, 57° to 106° F (13 8° to 41 1° C)
 Wiesbaden, 100° to 156° F (37 7° to 69° C)

Wildbad, 91.5° to 104° F (33° to 40°
C)

Table Waters

Apollinaris
Bellthal
Bilin
Birresborn
Cambrunnen
Condillac
Evian-les-Bains
Fachingen
Gellnau

Geiselstein
Giesshübel
Johannis
Krondorf
Kronthal
Malvern
Marcol
Renaissau
Rhens
Rörsdorf
Rosbach
Saint Galmier
Selters
Taunus
Thonau

SECTION A

THERAPEUTICAL CLASSIFICATION OF REMEDIES

Alteratives — Medicines which gradually change and correct a morbid condition of the organs, so that abnormal conditions become normal and metabolism is increased

Ammonium — Ammonii Chloridum

Antimony — Antimonii Oxidum, A Sulphuratum, A Tincturatum

Arsenic — Acidum Arsenicosum, Injectio Ferri Arsenitis, Liquor Arseniculus,

Liquor Arsenici Hydrochloricus, Liquor Sodii Arsenitis, Sodii Cacodylas

Calcium — Calcii Chloridum, Calcii Hypophosphis, Calcii Sulphidum

Iodine and the Iodides

Iron Salts

Mercury — Hydragric Cicta, Pilula Hydrarg, Hydrarg Perchloridum and

Sulchloridum, Hydrag Iodidum Rubrum

Phosphorus and the Hypophosphites

Potassium Salts

Sulphur — Precipitatum, Sublimatum and Sulphides

Tactible — Caffeinæ, Coca, Dulcamara, Guaiacum, Homidescimus, Mezereum,

Sassafrilla Sassafras, Tuxicum

Electics — Iridin, Leptandrin, Phytolaccin, Podophyllin

Anæsthetics — They are divided into *General* (by inhalation) and *Local* (by spray or other application to the part). **GENERAL ANÆSTHETICS** abolish consciousness and reflex action, and so prevent the perception of painful and other stimuli in the sensory centres. *Æther*, *Æther Methylatus* (sp gr 0.717), *Æthyl Bromidum*, *Æthyl Iodidum*, *A.C.E. Mixture*, *Carbon Tetrachloride*, *Chloroform*, *Chloroform*, *Kelene*, *Methylene*, *Narcotile*, *Nitrous Oxide Gas*, *Pental*, *Rignault's Anæsthetic Mixture*, *Somniform*. **LOCAL ANÆSTHETICS** prevent the reception of stimuli by the peripheral terminations of sensory nerves. *Acid Carbolic*, *Aconite*, *Æther* (spray), *Æther Methylatus* (sp gr 0.717), *Æther Methylicus*, *Æthoxyvitaminum*, *Æthyl Bromidum*, *Æthyl Chloridum*, *Alypin*, *Anæsthesin*, *Anæsthyll*, *Aromatic Oils*, *Benzoyl pseudotopine* (*Triopacocaine*), *Chloretone*, *Cocainæ Hydrochloridum*, *Cocainæ Phenylas*, *Coryl*, *Erythrophleinæ Hydrochloridum*, *Eucainæ Hydrochloride* (A) and (B), *Eugenol*, *Guaiacol*, *Guaiacyl*, *Holocainæ Hydrochloride*, *Ice*, *Iodoform*, *Kelene*, *Menthol*, *Methyl Chloridum*, *Methylal*, *Neivocaine*, *Nivamin*, *Novocaine*, *Orthoform*, *Orthoform New*, *Phenol*, *Camphor*, *Stovaine*, *Subcutin*, *Tropicocaine*, *Thymol*, *Yohimbine*

Analgesics or Anodynes — Medicines which alleviate pain by lessening the excitability of nerves or nerve centres. *Abiastol*, *Acetamidum*, *Acid Carbolic*, *Aconitum*, *Aconitina*, *Æthyl Chloridum*, *Agithin*, *Ammonol*, *Amyl Nitric*, *Antikuma*, *Antipyrine*, *Antiseptin*, *Antitoxic*, *Apocisin*, *Aristochin*, *Asaprol*, *Atropin*, *Belladonna*, *Bromides*, *Brucine*, *Butyl Chloral Hydras*, *Caffeinæ*, *Caryoputi* and *Caryophylli Ol*, *Camphor*, *Cannabis Indica*, *Chloral Hydras*, *Chloroform*, *Cimicifuga*, *Citrophen*, *Cocainæ Phenylas*, *Codeina*, *Codeinæ Iodas*, *Conina*, *Conium*, *Oreosotin*, *Dionine*, *Enquinnic*, *Exalgin*, *Gelsemium*, *Hyc cyanus*, *Ipecac Pulvis Compositus*, *Kryofin*, *Lactophenin*, *Lupulus*, *Malakin*, *Methylene Blue*, *Morphina*, *Opium*, *Orthoform*, *Papaver*, *Phenacetin*, *Phenalglin*, *Phenyl urethane* (*Euphorin*), *Piscidia*, *Pyrimidon*, *Quinna*, *Salipyrin*, *Sclorphen*, *Siloquinine*, *Scopola*, *Solanine*, *Spiritus Ætheris*, *Tolpyrin*, *Tolysal*, *Veratrina*

Anaphrodisiacs —Medicines which diminish the sexual passion Ammonii Bromidum, Belladonna, Camphora, Conium, Digitalis, Hyoscyamus, Lupulinum, Potassii Bromidum, Potassii Iodidum, Sodii Bromidum, Sodii Iodidum, Stramonium, Tabacum, also alkalis, hypnotics, depressants

Anhidrotics —Medicines which check perspiration Acid Acetic, Acid Camphoric, Acid Phosph Dil, A Salicylic, A Sulphuric Dilut, A Tannic, Agaricus, Agaricin, Atropina, Belladonna, Cotoin, Ergot, Ferri Sulphas, Ferri Mist Comp, Guaiacol, Camphorate, Mono- and Di-Camphorate, Scopola, Stramonium, Styrchnina, Zinci Oxidum

Antacids —Agents which reduce the acidity of the gastric contents Ammonia, Ammon Spirit Aromatic, A Carbonas, Bismuthi Trochiscus, Calen Hydric, C Carbonas Precipitatus, Calcis Liquor, C Saccharatus Liquor, Citra Preparata, Lithii Carbonas, Lithii Citras, Magnesia, M Carbonas, Potassa Liquor, P Carbonas, Pot Citras, Pot Tartas, Sapo Durus, Sodii Bic Phosphas

Mineral Waters —Contrexville, Ems, Fachingen, Tarasp, Vichy

Anthelmintics —Medicines which destroy intestinal worms (Vermicides), or expel them from the alimentary canal (Vermifuges)

VERMICIDES or Thread Worms —Acid Carbolic, Acon, Arecolinum Hydrobromide, Enema—Aceti, Eucalyptus Oil, Ferri Perchloridi, F Sulphatis, Olivæ Olei, Quassia, Ol Ricini and Terebinthinæ, Santoninum, Sodii Chloridi

Round Worm —Areca, Santoninum

Tape Worm —Acid Embelicum, Ammonii Embelic, Cusso, Embelia Rubes, Eucalyptus Oil, Extractum Filicis Liquidum, Guanati Cort, Kamala, Pelletierina Sulphas and Tannas, Terebinthinæ Oleum

VERMIFUGES Areca, Butea, Calomel, Cambogia, Jalapa, Kamala, Nucis Juglandis Spiritus, Ricini Oleum, Scammonium, Thymol Carbonate

Antidotes are mentioned under the several poisonous drugs

Antilithics —Medicines which counteract lithiasis or lithæmia, or tendency to the deposit of uric acid or urates, or to the formation of the corresponding Calculi Acid Nitric Dil, Acid Phosph Dil, Cystamine, Hexamethylureteramine, Lithii salts (see p 733), Magnesi Mag Carbonas, Magnesi Citras, Piperazine, Piperazine Quinate, Piperidine Tartate Acid, Podophyllum, Potassii Acetas, Potassii Bicarb, Sapo Durus, Soda Tartarata, Sodii Citras, Sodii Citro-tart Effor, Sodii Phosphas, Urea, Uresii

diuretics, saline purgatives

Mineral Waters —Carlsbad, Friedrichshall, Hunyadi Janos, Meigenthheim, Neuenahr, Selters, Teplitz, Vals, Vichy, Wildungen

Antiperiodics —Medicines which have the property of preventing the periodical attacks of certain fevers Acid Arsenios, Beberina Sulph, Berberis, Cinchona, Cinchonidine Salicylate, Cusparia, Ferri Arsenio-Citras Ammonata, Ferri Aisenatis Inj, Nalcotina, Nectandra Cort, Quinine Salts, Picrorhiza, Salicin

Antipyretics —Medicines which reduce and control the temperature in fever Acetopyrin, Acids, Acid Acetyl salicylic, Acid Anisic, Acid Salicylic, Aconitum, Ammon Benz, Ammonii Salicylas, Ammonol, Ammonol Bromide, Lithiated and Salicylate, Antibacillus, Antipyrine Camphorate, Antiseptin, Antitoxine, Antimonium Tartaratum, Apolysin, Aristochin, Asaprol, Aspirin, Bromopyrin, Camphor Salicylate, Chinaphenon, Citophen, Eupyrine, Kiyofin, Lactophenon, Malakin, Magnesi, acid, Paraphenetidin Agarate and Camphor, Phenazonum, Phenocoll Hydrochloridum, Phesin, Piperina, Potassii Citras, Pyramidon, Pyramidon Mono- and Di-Camphorate and Salicylate, Pyrantin, Pyridin, Quibricin, Quinina, Resorcin, Salicin, Antis, Sodii Di-Thio-salicyla, Sodii Sâhevlas Sodii Sulphocarbolas, Spirit Æther Muraticus, Spirit Æther Nitrosi, Spirit Rectificatus, Thallina Sulphas, Tolpyrin, Tolsal, Triphenine

Antiseptics — Agents which prevent decomposition by inhibiting the growth of Micro organisms. Acid Benzoicum, Acid Boricum, Acid Carbolicum, Acid Chromicum, Acid Cinnamic, Acid Cresotinum, Acid Cresylicum, Acid Hydrochloricum, Acid Nitricum, Acid Oxynaphthoicum, Acid Pyrogallicum, Acid Pyroligneosum Crudum, Acid Salicylicum, Acid Sulphocarbolicum, Acid Sulphurosum, Acid Trichloroaceticum, Albargin, Aluminium Acetatis Liquor, Alum Aceto Tartaric, Actol, Alum Chloridi Liquor, A Oleas, A Nitras, A Sulphas, Alumnol, Ammon Benzoas, Amylotoim, Antiseptin, Antinosin, Anytin, Anytol, Argentamin, Argentol, Argonin, Aristol, Aseptin, Asterol, Balsamum Peruvianum and Tolutanum, Benzoin, Benzonaphthol, Benzoyl Peroxide, Betol, Bismone, Bismuthi Benzoas, B Betanaphtholus, B Cera Salicylas, B Cinchonidine Iodidum, B Diethio salicylas, B Iodoresorcin Sulphonis, B Oleas, B Oxychloridum, B Oxyiodogallas, B Phenol, B Phosphas, B Quinolme Sulphocyanidum, L Salicylas, B Subgillus, B Subiodas, B Sulphas, L Tribromphenolus, Borax, Boro Glycende, Bimulcan, Calci Iodid, Calci Purificatum, Cukinol, Cuka Chlorinata, Carbo Ligni, Camphor, Camphor Phenol, C Resorcin, C Salicylate, C Thymol, Carbonis Bisulphidum, Caryophyllum, Chmoline, Chmoline Tetrat, Chmosol, Chlori Liquor, Chloroform, Cinchonum et Iodo Sulphas, Cinchonidine Sulphocyanobolas, Cinnamomum, Cinnamomi Ol, Collargol, Copaiba, Cresotum, Cupri Oleas, C Sulphocarbolus, Cyllin, Cysamine, Dextroform, Diaphtherin, Diaphthol, Diiodoform, Eau de Javelle, a Eigon, Eka Iodoform, Eucalyptol, Eucalyptus Oil, Eudoxin, Eugallol, Euguform, Eupyrin, Euprophon, Fel Bovinum Purif, Fluoroform, Formaldehyde, Formicin, Fortom, Glaciline, Glutol, Glycerinum, Glycosal, Guaiacol and its salts, Guaiacuin, Guaiamar, Guaiasanol, Helenn, Hermophenyl, Hexamethylene tetramine, Hydrargyri Cyanidum, H Ethylene diamine Citras, H et Potassu Iodidum, H Nitratis Liquor Acidus, H Nuclemas, H Perchloridum, H Salicylas, H Subchloridum, Hydrag Zincu Cyanid, Hydragyrol, Hydrogeni Peroxidi Liquor, Hydronaphthol, Ichthyrgan, Ichthoform, Iodi Tribromidum, Iodi Trichloridum, Iodofan, Iodoformin, Iodoformogen, Iodoformum, Iodol, Iodolene, Iodopyrin, Iodum, Irol, L'al Lactonaphthol, Longillol, Listerine, Loretin, Losophan, Lysoform, Lycol, Menthol, Menthosol, Menthoxol, Mercuamine, Metakalin, Methyl Salicylas, Microcidine, Naphthalene, Naphthol, Naphthol Camphor, Nosophen, Nuclein and its salts, Orthoform, Para monochlorophenol, Phenosalyl, Potassa Sulphurata, Potassu Permanganas, Protargol, Pyoktanin, Quinaphthol, Quinine Hydrochlor, Quinina Sulphas, Resorcin, Resorcinol, Sal Alembroth, Saligallol, Salitannol, Salol, saloquinine, Sypiol, Sanoform, Soda Chlorinate Liquor, Sodii Anis, S Benzoas, S Chloridum, Sodii Diethio salicylas, S Fluoridum, S Salicylas, S Silicofluoridum, S Sulphas, S Sulphocarbolus, Solveol, Soziodol, Strontu Salicylas, Sulphammol, Tachiol, Teiebinthum et Oleum, Thallema Sulphas, Thymol, Traumatol, Tribromphenol, Tribromoresorcin, Trichlorphenol, Trikresol, Vioform, Yeast, Zinci Chloridum, Zinci Sulphas, Zinci Sulphocarbolus, Zymocido

Antispasmodics — Medicines which allay or prevent the recurrence of spasms. Acid Hydrocyanic Dil, Aether, Aether Aceticus, Aethyl Iodidum, Ammonia Liquor, A Carbonis and Bicarbonis, Spiritus Ammonici Aromaticus, Ammoniacum, Amyl Nitris, Amyl Valerianus, Intim Tartaratum, Argenti Nitras, Argenti Oxidum, Asufetida, Atropine Valerianus, Belladonna, Boldo, Bromides, Cajuput Ol, Calendula, Camphora, Camphora Monobromata, Cannabis Indica, Caryophyllum, Cistoreum, Cera Ovala, Chloral Hydras, Chloroformum, Cimicifuga, Conium, Ethyl Nitritus Liquor, Euphorbia Pululifeia, Galbanum, Grindelia, Hyoscyamus, Iso butyl Nitrite, Juniperi Ol, Lobelia, Menth Pip Ol, Moschus, Opium, Physostigma, Physostigmine Salicylas and Sulphas, Pil Aloes et Asafetide, Piscidia, Quebracho, Ruta Oleum, Santonin, Sodii Nitris, Spir Ammon Fetid, Stramonium, Sumbul, Tabacum, Teiebinthma, Trinitrin Liquor and Tabella, Valeriana and Valerianates, Zinci Oxidum, Zinci Sulphas, Zinci Valerianas

Aperients — See Cathartics

Aphrodisiacs — Medicines which increase sexual appetite. Alcohol, Belladonna, Calcu Hypophosphis, Camphor, Cantharis, Coffee, Damiana, Tinct Ferri Perchlor, Hematinics and Nux Vomica (strychnina), Phosphorus, Tonics, Yohimbin

Aromatics.—*See* Cumminatives

Astringents.—Medicines which produce contraction of the tissues, diminution in the size of blood-vessels and coagulation of the albuminous fluids, they are given to improve digestion and check secretions, mucous discharges, and hæmorrhages, or applied topically to stop bleeding and diminish discharges.

Mineral Substances.—All the Diluted Mineral Acids, Aluminium Salts, Argentamin, Argenti Nitras, A Oxidum, Bismuth salts, Borax, Cadmi-
na, Calci Carbonas Præcip, Calci Hydrias, Cupbolic
Creta Præp, Cupri Sulphas, Ferri Perchlor. Liquor,
F Pernit Liquor, F Sulphas, Ferri et Quin Cit, Plumbi Acetas,
P Carbonas, P Oxidum, P Subacetatis Liquor Fortis, Zinci Acetas,
Z Carbonas, Z Chloridum, Z Oxidum, Z Sulphas, Z Sulphocarbolas

Vegetable Substances.—Acetum, Acid Acetic Dil, A Gallic, A Tannic,
Aieca, Belle Confectio, Catechu, Cinchona, Cinnamomum, Coto, Ergoti,
Erigerontis Oleum, Filix Mas, Galla, Gallogen, Glutanol, Granati Cort,
Guarana, Gummi Eucalypt, Hamatoxylum, Hamamelis, Honthin,
Hydiastis, Ispaghula, Krameria, Kino, Lani, Matico,
Juglandis Spiritus, Opium, Quercus, Rheum, Ros-
Symphytum, Tannalbin, Tannigen, Tannoform, Tanocol, Tunnone, Tere-
binthina Ol, Ulmus, Uva Uris, Vinca Major

Carminatives.—Medicines which stimulate or aid the removal of flatus from the stomach and intestines, and relieve griping. Æther, Æther Aceticus, Anethi Ol, Anisi Ol, Asafetida, Boldo, Camphor, Carbo Ligni, Cardamomum, Carii Ol, Caryoph, Cascarella, Chloroformum, Cinnamomum, Coriander, Coto, Fœniculum, Ipecacuanha, Jumper, Lavand Ol, Limon Ol, Menth Pip Ol, Menth Virid Ol, Menthol, Menthol Valerianate, Myristica, Myrtila, Pimento, Piper, Rosmarini Oleum, Sumbul, Valeriana, Zingiber

Cathartics.—Medicines which promote intestinal evacuations

Apertive or Laxative.—Amygdalæ Oleum, A Hydrochloridum,
Asafetida, Baptisin, Belladonna, Cascara - Pulpa, Enoani-
min, Fel Bovinum, Ficus, Glycyrrh Pulv Comp, Ipecac Juglans,
Lactuca, Lani Oleum, Magnesia, M Carbonas, M Citratis Liquor,
Manna, Mel Depuratum, Menyanthes, Mori Succus, Nux Vomica,
Olive Oleum, Phenolphthalein, Potassu Sulphas, P Tartias, Plinum,
Rhamnus, Ricini Oleum, Sapo Durus, Sodii Citio-tartias
Sodii Phosphas, Sodii Sulphas, Sodii Sulphovinas, Soda tartarata,
Sulphur, Sulphur Præcip, Tamarindus, Tanaxacum

Purgative.—Acid Cathartic, Aloes Barb, A Socot, Alom, Baptisin, Col-
chicum, Convallarin, Helleborus Niger, Hydiarg Subchloridum, Iri-
septandum, Magnes Sulphas, Mangan Sulphas,
Purgatin, Rheum, Senna, Sodii Chloridum,

Diastic or Hydragogue.—Apocynum, Bryonia, Cambogi, Colocynthis,
Crotonis Oleum, Elaterum, Elaterium, Helleborus Niger, Hydiarg
Creta, Hydiarg Subchloridum, Jalapa, Kaladana, Lobeia, Magnes
Sulphas, Potass Tart Acidus, Scammonium, Sodii Sulphas, Veratrina

Mineral Waters.—Achseimannstern, Birmenstorf, Carlsbad, Friedrichshall,
Homburg, Hunyadi-Janos, Kissingen, Marienbad, Pullna, Royal Hun-
garian Bitter Water (Buda-Pesth), Seidlitz

Caustics.—Medicines which destroy the vitality of the parts to which they are
A Aisemosum, A Carbolicum, A Chloromicum,
A Nitricum, A Pyrogallic, A Sulphurici Pasta, A Trichlor-
acetic, Alum, Acetum, Acetum Aceto-Tart, Antum Chloridi Liquor,
Argenti Nitra- Cupri Acetas, C Nitras, C Subacetas,
C Sulphas, Ferri Pernitratiss Liquor, Formaldehyde, Hydi Iod Rubr, Hydr
Nitratis Acidus Liquor, Hydr Ox Rubr, Hydi Ox Flav, Hydr P
Potassa Caustica, Potassa c Calce, Potassu Permang, Sodii Eth
Zinci Chloridum, Zinci Nitras

Cholagogues—The *direct* increase the amount of bile secreted. Acid Hydrochlor Dil, Acid Nitric Dil, Acid Nitrohydrochlor Dil, Aloes Ammonii Chloridum, Ammonii Phosphas, Antim Sulphuratum, Boldo Colchicum, Colocynth, Eucatalol, Eunosmus, Fel Bovinum Purif, Hydrastis, Ipecacuanha, Iridin, Jalapa, Phytolaccin, Podophyllum, Rheum, Sodii Benzoas, Glycocholas, Salicys, and Sulphur. The *indirect* act by stimulating the duodenum. Mercury, especially the Subchloride most Cathartics.

Mineral Waters—Ems, Friedrichshall, Hungarian, Hunyadi Janos, Kis-singen.

Counter-Irritants—Substances which stimulate and cause irritation or inflammation of the parts to which they are applied, they differ in their intensity of action, and may be divided as follows:—

RUBRIFICANTS—Agents which when applied to the skin, produce local warmth and redness. Acid Aceticum, Aether, Alcohol, Ammonia Liquor, Ammonium, Ammonia, Emp Cicutarum, Cantharis, Iodum, Lin Camphorae Ammon, Lin Capsici, Lin Chloroformi, Lin Sinapis, Lin Iodi Fortis, Mentholi Emp, Mezerium Ol Cadinum, Ol Cajuputi, Ol Limonis, Ol Rosmarini, Ol Rutae, Ol Succini, Ol Terebinthi, Picis Emp, Thymol, Ung Ictoni.

VESICANTS OR ECTHASICS—Those which raise a vesicle or blister. Acidum Aceticum Glaciale, Ammonia Liquor Fortior, Cantharidin, Cantharis Emp, Ecthasicus Liquor, Euphorbium, Mezerium Ung, Rutae Oleum, Sinapis Lin, Sinapis Oleum.

PUSTULANTS—Those which produce pustules. Antimonium Tartaratum, Argentum Nitras, Crotonis Oleum.

Demulcents—Substances which protect, and thus allay irritation of, the mucous membranes. Acacia Gum, Althaea, Amygdali Dulc and Oleum, Amylum, Canagren Cetraria, Cydonium, Cynoglossum, Ficus, Gelatinum, Glycerinum Boracis, Glycyrrhiza, Hordeum, Ichthyocolla, Ispaghula, Linum Maianthi, Mel Depuratum, Oliva Oleum, Ovi Albumen, Pinnum, Sacch Purificatum, Salep, Sovum, Theriaca, Thymocantha, Triticum Repens, Ulmus, Uvae.

Deodorants—Substances which destroy offensive odours and absorb foul gases. Chlorine and its oxides, Acid Chromic, Acid Nitric, Acid Sulphurous, Bromum, Calx, Calci Permanganas, Calbo Ligni, Chinosol, Diiodoform, Eucalypti (A) and (B), Fla iodoform, Eucalypti Ol, Europhen, Ferri Sulph, Formaldehyde, Hydrogeni Peroxidum, Iodoformin, Iodoformogen, Iodoformum, Iodol, Iodolene, Iodum, Loretin, Mentholol, Naphthalene, Nosophen, Paraformaldehyde, Plumbi Nitrates, Potass Permang, Resorcinol, Thymol, Trichlorophenol, Vioform, Zinci Chloridum.

Depilatories—Chemicals which destroy living hair. Barri Sulphid, Calx Sulphurata, X Rays.

Desiccants—Agents which check secretion, and allay discharges from ulcers and wounds. Acidi Borici Pulvis, Bismuthi Subnit, Calamina, Calci Carbonas Princip, Calci Hydras, Cieta Prapuata, Magnesi Carbonas, Plumbi Acetas, Plumbi Carbonas, Tale, Zinci Carbonas and Oxidum.

Diaphoretics—Medicines which increase the action of the skin and induce perspiration. Acidum Salicylicum, Aconitum, Aether, Alcohol, Ammonia Acetatis Liquor, Ammon Carbonas, Ammon Chlorid, Ammon Citratis Liquor, Ammon Phosphas, Antimonialis Pulvis, Antim Vinum, Antim Sulphurata, Aconitine Hydrobrom, Amomacia, Buchu, Cajuputi Sp and Oleum, Calendula, Camphor, Chinosol, Chloroform, Colchici Vin, Doveri Pulv, Dulcamara, Eupatorium, Grindelia, Guarici Ammon Tinct, Ipecac Pulv Comp, Ipecac Vin, Jaborandi, Lactuca, Lobelia, Mezerium, Morphina, Opium, Pilocarpina, Pilocarpine Hydrochloride, Nitrate and Salicylate, Potass Acetas, Potass Citratis, Potass Nitrates, Silium, Sassafras, Senega, Serpentina, Sumatuba, Sodii Salicylas, Sp Aethoris Nit, Spiritus Camphorae, Spiritus Rectificatus, Sulphur, Sulphur Princip, Terebinthini Oleum.

Disinfectants—Substances which destroy the specific microbes or toxins of communicable diseases. Acid Cubol, Acid Chromic, Acid Cresylic, Acid Nitrosum, Acid Pyrogallie, Acid Sulphurousum, Aluminium Chloride Liquor,

Glycyrrhiza, Guaiacul and its salts, Iodides, Ipecacuanha, Lanius Cortex, Lobelia, Myrica, Physostigma, Pinu Oleum, Pix Liquida, Quillai, Sulla, Senega, Styrae Pæp, Sulphur, Terpene Hydrate, Terebene, Terebinthi Oleum, Vapores Acidi Carbonici, Chlori, Creosoti, and Iodi, Verba Santa

Febrifuges — *See* Antipyretics

Galactagogues — Agents which increase the secretion of the mammary gland
Jaborandi, Potass Chlorat, and Tonics

Hæmatinics — *See* Tonics, Blood

Hæmostatics — *See* Styptics

Hypnotics — (Soporifics) — Medicines which induce sleep, and thus remove the consciousness of pain by lessening the excitability and functional activity of the brain cells. Acetophenone, Acid Hydrobrom Dil, Ammon Bromidum, Amylene Hydrate, Antispasmod, Apomorph Hydrochlor, Boldo, Bromural, Camphor, Camphora Monobromata, Cannabis Tenuis, Cannabis Ind, Chloral Hydris, Chloralhydrate, Chloralose, Chloroform, Chlorobrom, Codina, Conium, Dormiol, Dromine, Eucodal, Heroin, Heroin Hydrochloride, Hyoscyamus, Hyoscin, Hyoscine Hydrobromide, Hydrochloride and Hydrodide, Hypnol, Hypnone, Lactophenol, Lithi Bromidum, Lupulus, Metaldohyd, Methylal, Morphin, Morphin Bimeconatis Liquor, Nuxcina, Nemonal, Opium, Papaver, Papavina, Paraldehyde, Peronine, Piscidia, Potass Bromidum, Scopolamine Hydrobrom, Sodii Bromidum, Somnol, Stramonium, Sulphural, Tonal, Trional, Urethane, Veronal

Laxatives — *See* Cathartics

Mydriatics — Drugs which produce dilatation of the pupil. Atroline Hydrobromide, Atropin, Atrop Methylbrom, Atropine Salicylate, Atroscine, Belladonna, Cocaine, Cocaine Hydrochloridum, Datura, Duboisina Sulphas, Ephedrine Hydrochloride, Euphrasine Hydrochloride and Salicylate, Homatropin, Homatropine Hydrobromidum, Hydrochloridum and Salicylis Hyoscyamus, Hyoscin Hydrobromidum and Hydrochloridum, Mydin, Mydinum, Myoscymin Hydrobromas and Sulphas, Scopol, Stramonium

Myotics — Drugs which contract the pupil. Esarine, Jaborandi, Morphin, Opium, Physostigma, Physostigmine Salicylas and Sulphas, Pilocarpina

Narcotics — *See* Hypnotics

Nutritives — Substances which aid assimilation and improve the condition of the tissues. Acorus Gum, Amygdala Dulc, Byns Extractum, Calci Glycophosphas, Camu Extract, Carageen, Cetarii Decoctum, Ficus, Foidum, Manna, Maranta, Mel Depuratum, Morilla Ol, Olive Oleum, Prunum, Sacchi Lactis, Sacchi Purificatum, Salep, Sevum, Somatose (various), Sp Vini Gallici Mist

Parasitocides — Medicines which destroy vegetable and animal parasites. Acid Cubic, Acid Pyrogall, Acid Salicylic, Acid Sulphurosum, Anthiarobin, Chrysarobinum, Cupri Oleas, Hydi Nitrat Ung, Hydi Oleas, Hydi Oxid Itub Ung, Hydi Perchloridum, Iodi Pigmentum, Mercurial preparations, Olea Expressa et Essent, Naphthylene, Picrotoxin, Potassa Sulphurata, Pyrethrum Roseum, Quassia, Sozoiodol, Staphisagria, Stylicus Unguentum, Sulphur, Tabacum, Thymol

Purgatives — *See* Cathartics

Pustulants — *See* Counter-Irritants

Refrigerants — Agents which relieve febrile thirst, and impart a feeling of coolness. Actum, Acidum Aceticum, A Citricum, A Hydrochlor Dil, A Nitric Dil, A Phosph Dil, A Sulph Dil, A Tartaricum, Ammon Acet Liquor, Aqua, Auranti Succus, Imperial Drink, Limonis Succus, Magnesi Citratis Liquor, Mori Syrup, Oxymel, Potass Citras, Potass Chloras, Potass Nitrates, Potass Tart Acid, Prunum, Sp Ether Nitr, Sp Ether Muriaticus, Tamarindus. *See also* Diaphoretics and Antipyretics

Rubefacients — *See* Counter-Irritants

Sedatives—Medicines which exert a soothing influence, by diminishing pain, depressing vital activity, or tranquillising abnormal muscular movement.

Local—Acid Carbolic, Acid Hydrocyan Dil, Atropina, Belladonna, Borax, Chloral, Creosotum, Morphine, Opium, Plumbi Acetas, P Carbonas, P Subacetatis Liquor Dilutus—See also Anesthetics (Local), and Anodynes.

Respiratory—Acid Hydrocyan Dil, Ether, Ether Aceticus, Ethyl Iodidum, Ammon Bromid, Amyl Nitrite, Belladonna, Cannabis Indica, Chloroformum, Chloral, Codonv, Codonv Hydrochloride, Phosphate and Sulcyate, Conium, Conium, Conium Hydrobrom, Conium Hydrochloridum, Dione, Gelsemium, Heroin, Heroin Hydrochloride, Hyoscyamus, Lactucarium, Lantocerasi Aqua, Lobelia, Morphine and salts, Nuxal, Opium, Potassum, Prun Virgin Syrup et Tinct, Stramonium, Terebinthine Oleum.

Nervine—Acid Hydrobrom Dilutum, Ammon Bromidum, Animon Valerianas, Amyl Valerianas, Antum Tartaratum, Antispasmod, B-Ergon, Bromethylformine, Camphora, Camphor, Gallobromol, Gelsemium, Hyoscyamine Hydrobromidum and Sulphur, Lith Bromidum, Lupulum, Lupulus, Liquor, Menthol Valerianate, Nemonia, Nucum Bromianum, Sassa, Phenacetin, Phenazonum, Physostigma, Piscidia, Potass Bromidum, Salix Nigra, Scutellaria, Soda Bromidum, Thionil, Valyl, Veratrum Viride, Veronal, Viburnum, Zinc Bromidum.

Gastric—Acid Arsenosum, Acid Carbolic, Acid Carbonic, Acid Hydrocyan Dil, Acid Phosph Dil, Ammon Bromidum, Argenti Nitras, Argenti Oxidum, Belladonna, Bismuth salts, Calcii Hydrias, Calcis Liquor, Ceru Oxalas, Chloral, Chlorobrom, Chloroform, Cocaine Hydrochlor, Cocaine Phenylas, Creosotum, Hydragric Creta, Hyd Subchlor (small doses), Hyoscyamus, Ice, Ipecacuanha (small doses), Opium, Papaver, Phloidylin, Potass Bicarb, Potass Bichrom, Potass Bromid, Soda Bicarb, Soda Bromidum, Zinc Oxidum.

Cardiac—Acidum Hydrocyanicum Dilutum, Aconitum, Amyl Nitris, Antum Tart, Apocynum, Aqua Lantocerasi, Belladonna, Conium, Chloral, Digitalis, Ergota, Hyoscyamus, glycerinum, Opium, Scilla, Soda Nitris, Spirit Aether, Veratrum Viride.

Sialagogues—Medicines that increase the secretion of the saliva. Acetum, Acid Acetic, Acid Citric, Acid Tartaric, Aether, Alcohol, Aescoline Hydrobromide, Armoacia, Amantium, Dilute Acids, and Acid salts, most Emetics (especially Antimony and Iodine), Hydrogynum and its salts, Iodides, Ipecacuanha, Jaborandi, Limonis Succus, Mezerium, Physostigma, Pilocarpine, Pilocarpine Hydrochloridum, Nitras and Salicylis, Piper, Pyrethrum, Rheum, Sassa, Tabacum, Tamarindus, Zingiber.

Soporifics—See Hypnotics.

Sternutatories—Medicines which cause sneezing, and increase the nasal mucous secretion. Ipecacuanha (powdered), Tabacum (snuff), Veratrum Viride (powdered).

Stimulants—Medicines which increase the function of a part, or of an organ.

Cerebral—Absinthum, Caffein, Theobromine.

Nervine—Acid Arsenosum, Aether, Ammon Arom Spt, Ammon Carb, Ammon Chlorid, Ammon Phosph, Asafetida, Belladonna, Calcii Hypophosphis, Cannabis Ind, Cinthius, Castoreum, Cocca, Coffee, Ergota, Guarana, Hydrastis, Kola, Musk, Nux Vomica, Oleum Cuputi, Phosphorus, Spirit Ammon Felid, Stychina, Valeriana.

Stomachic—See Carminatives and Stomachic Tonics.

Circulatory—Aether, Aether Aceticus, Aether Spiritus Nitrosi, Alcohol, Ammonia Aromat Spt, Camphor, Convallaria, Digitalis, Stychina, Sumbul.

Local—Potass Chloras. See also Counter Irritants.

Stomachics—Medicines which directly promote the functions of the stomach and improve the appetite and digestion. See Carminatives, and Tonics, Stomachic.

Styptics—Remedies which arrest bleeding—Aceticum, Acid Sulphuric Dil, Acid Tannic Adrenalin, Albumen, Aluminium Oleate, Argenti Nitrate, Benzoin, Bryonia, Catechu, Chinisol, Cinchona Pulvis, Colloidium Calci Chloridum, Cornutine Citrate, Cornutine Hydrochloride, Cotinine Phthalate (Styptol), Creosote, Cupri Sulphas, Cupri Sulphocarbols, Eigota, Ergotinine, Eriogerontis Oleum, Ferri Perchlor Liquor and other Ferric salts Ferri et Ammonii Sulphas, Ferripyrin, Gallæ, Glycerati Cort, Gummi Rubri Extractum Liquidum, Hamamelis, Hamamelis, Hydrastis, Hydrastinis Hydrochloridum, Kino, Krameria, Matico, Opium, Plumbi Acetas, Plumbi Subacetatis Liquor, Quercus, Quinina et Ferri Chloridum, Quinine Hydrochlor, Salipyrin, Spiritus Rectificatus, Stypticum, Suprarenal Gland and Extract, Terbinthinæ Oleum, Zinci Acetas, Zinci Sulph

Sudorifics—*See* Diaphoretics—When diaphoretics act very powerfully, they are called sudorifics

Tonics—Therapeutic agents which impart strength or tone to the functions of the body or its parts

Acting through the blood and improving its qualities—Acid Arsenosum, Acid Phosph Dil, Acidum Nucleum and salts, Alboferin, Carminferin, Foston's Syrup, Ferri Acetatis Liquor, Ferri et Ferri, Ferri Albuminas, Ferri Alginas, Ferri Arsenas, Ferri Bromidum, Ferri Cacodylas, Ferri Saccharatus, Ferri Ammonii Citras, Ferri et Quinina Citras, Ferri Chloroxydum, Ferri Hypophosphis, Ferri Iodidum, Ferri Liquor Dialysat, Ferri Lactas, Ferri Oxid Magnet, Ferri Perchlor, Ferri Pernit Liquor, Ferri Peptonas, Ferri et Man Peptonatis Liq, Ferri Phosphas, Ferri Phosph Co Syrup (Squire), Ferri Pilula, Ferri Redactum, Ferri Sulphas, Ferri Tartaratum, Ferripyrin, Ferriethol, Glycerophosphates, Hæmoglobin and preparations, Lecithogen, Morrhu Oleum, Potass Permang, Sodii Cacodylas, Sodii Methylarsenas, Syri Calci Lactophos et c Ferri, Syrup Hypophosph Co

Nervine—Acid Arsenosum, Argenti Nitras, Argenti Oxidum, Calci Hypophosphis, Cinchona, Coca, Cupri Sulphas, Damanas, Ferrum salts, Glycerophosphates, Guarana, Lecithin, Morrhu Oleum, Nux Vomica, Phosphorus, Quinina, Sodii Hypophosphis, Strychnina, Sumbul, Zinci Acetas, Zinci Oxidum, Zinci Phosphid, Zinci Sulph, Zinci Valerianas

Stomachic and Intestinal—Acid Hydrochlor Dil, A Nitric Dil, A Nitrohydrochlor Dil, A Phosph Dil, A Sulph Dil, Aloes, Anthemis, Armoracia, Aurant Cort, Bebeerina, Bebeerinæ Hydrochlor and Sulphas, Berberis, Boldo, Buchu, Calumba, Canellæ Cortex, Capsicum, Cascarella, Chiretta, Cimicifuga, Cinchona, Cinchonidinæ Salicylas, Cinchonidina, Cinchonina, Cusparia, Decoct Aloes Comp, Eriogerontis Oleum, Eupatorium, Gentiana, Guarana, Hydrastis, Ignatia, Krameria, Kava Kava, Limonis Cortex, Lupulinum, Lupulus, Menyanthes, Mezereum, Nectandra, Nux Vomica, Orexin, Orexin Hydrochloride, Orexin Tannate, Pancreatic Enzymes, Pareira, Pepsin, Peptonised Foods, Piper, Quassia, Quebracho, Quinina Hydrochloridum, Quinine Sulph, Rheum, Salicin, Sarsaparilla, Serpentina, Simaruba, Sinapis, Sodii Chloridum, Strychnina, Taraxacum, Ulmus, Uva Ursi

Cardiac—Acidum Arsenosum, Adonis, Adrenalin, Adrenalin Chloride, Ethoxycaffeinum, Apocynum, Caffeina, Caffeina Sodio Benzoas, Caff Sodii Salicylas, Convallaria, Convallamarin, Digitalis, Digitalein, Digitalin, Digitin, Digitonin, Digitoxin, Diuretin, Erythrophleum, Erythrophleum, and Hydrochloride, Ferum salts, Nux Vomica, Oxysparteine, Oxysparteine Hydrochloride and Sulphate, Scilla, Sparteine Sulphas, Strophanthus, Strychnina, Suprarenal Gland, Veratrum Viride

Mineral Waters—Adelheidsquelle, Alot, Altwasser, Auteuil, Berka, Bocklet, Gasten, Kreuznach, Membeigh, Orezza, Pyrmont, St Moritz, Spa, Schwalbach, Wildungen

Vaso-Dilators—Amyl Nitrite, Erythrol Tetranitrate, Mannitol Hexanitrate, Sodii Nitras, Trinitrin

Vermicides and Vermifuges—*See* Anthelmintics

Vesicants—*See* Counter Irritants

SECTION B

REMEDIES EMPLOYED IN SPECIAL AILMENTS

Abortion, Threatened Ergot in small doses, Hydrastis, Morphina, Opium, Plumb. c. Opio Pil, Potass. Chloras, Viburnum Prunifolium

Abscess (to abort) *Internally* Aconite, Belladonna, Sulphides *Locally* Chloro Liqueor, Glycerinum Belladonnae, Iodoformum, Iodum, Acid Boric, A Carbolicum, Argenti Nitras, Menthoxol, Pot. Permang

Acne *Topically* id Nucleic, Liqueor Arsenicals, Calci Sulphidum, Vinum Ferri Cit, Levurine, Saline Purgatives, Confect. S. B. L., Yeast *Locally* Belladonna, Benzoin, Calamine, Hydrag. Perchlor. (Lotio), Ichthyol, Potassa Sulphurata, Lotio Zinc. Oxid, Resorcin, Sulphur, Ung. Sulphuris Hypochlor., Ung. Sulphuris Iodid

Aque—See *Fever, Malarial*

Albuminuria Ammon. Acet. Liq., Antim. Tart., Digitalis, Ferri Perchlor. Tinct., Jalapind., P. Jalapae Co., Milk, Nitroglycerin, Potass. Acet., P. Bitart., P. Bicarb., P. Citras., P. Iodid., Saline Purgatives, Sodii Nitras, Strontii Lactas, Supra-renal Gland

Alcoholism Ammoniae Acetat. Liqueor, Ammon. Bromid., Ammon. Carb., Arnica, Arsenic, Calumba, Capsicum, Cimicifuga, Cinchona, Cocaine, Hydrochlor., Gentiana, Hyoscin. e Hydrobrom., Lupulus, Nux. Vomica, Opium, Quinina, Atropine and Strychnine—See also *Delirium Tremens*

Alopecia Areata Acetum Cantharidis, Acid Lactic, Acid Salicylic, Chrysarobin Ung., Hydrag. Perchlor. (Lotio), Liniment Ammon., Lin. Camph. Ammon., Lin. Chloroform, Linimentum Cinale, Lin. Crotonis, Lin. Sinapis, Lotio Cinalis, Lotio Stimulans, Pilocarpine Nitras, Sulphur

Alteratives—Section A

Amenorrhoea Aloes, Apoc. Capsules, Auri et Sodii Chloridum, Calendula, Cantharis, Cimicifuga, Ergota, Ferri Bromidi Syrupus, Ferri Carb. Sacch., Ferri Lactas, Ferri Phosphas, Ferrum Redactum, Guaiaci Resina, Mistura Ferri Co., Menyanthes, Myrica, Pil. Aloes et Myrica, Potass. Permang., Ruta Oleum, Saline Purgatives, Sinapis, Hamamelis, Nervine Tonics

Anæmia—See *Tonics, Blood*, Section A

Anæmia—Pernicious Acid Arsenosum, Acid Nucleicum and salts, Acid Salicylic, Bone Marrow, Ferri Glycero-phosph., Hæmoglobin and preparations, Hydrag. c. Creta, Hyd. Subchlor., Phosphorus, Strychnina, oral, gastric and intestinal antiseptics, Anti-streptococcus serum

Anæsthetics—Section A**Analgesics or Anodynes**—Section A

Anasarca—See *Dropsy*

Aneurysm Aconitum, Amyl Nitris, Gelatin injected subcutaneously, Morphina, Potassu Iodidum in very large doses, Strontii Iodid

Angina Pectoris Acid Arseniosum, Acid Hydrocyanic Dil, Æther, Ætheris Spt, Ætheris Nitrosi Spt, Æthyl Bromidum, Alcohol, Ammon Arom Spt, Amyl Nitris, Argenti Nitras, Belladonna, Chloralamid, Chlorof Spt, Erythiol Tetranitrate, Ethyl Nitritis Liq, Iso butyl Nitris, Morphina (hypoderm), Nitroglycerin, Potass Iodidum, Pyridin, Sodii Nitris

Anhidrotics —Section A

Inkylotomiasis Chloroform, Eucalypt Ol, Filicis Maris Ext Liq, Saline Aperients, Thymol

Antacids —Section A

Anthelmintics —Section A

Anthrax Acid Carbolic (injection), Anti anthrax serum, Chloride of Zinc Points, Potass Caustic

Antidotes —Section A

Antilithics —Section A

Antiperiodics —Section A

Antipyretics —Section A

Antiseptics —Section A

Antispasmodics —Section A

Aperients —Section A

Aphrodisiacs —Section A

Aplthæ Acidum Boricum, Acidum Sulphuricum, Alum (pulv), Argenti Nitras, Glycerinum or Mel Boracis, Myrrha, Potass Chloras

Opophry Aloes, Cloton Ol, Melitenum, Hydrarg Subchlor, Pulvis Jalape Co, Ricini Ol, Terebinth Encma, Stimulants contra indicat

Aromatics —Section A

Arterio sclerosis Erythiol Tetranitrate, Potass Iodid, Sodii Nitris, Sodii Salicyl Digitals and other circulatory stimulants to be avoided

Ascarides —See *Anthelmintics*

Ascites —See *Dropsy*

Asthma Acidum Arsenosum, Acid Hydrocyanicum Dilutum, Æther, Æthyl Iodidum, Ammon Fetid Spirit, Ammoniacum, Ammonii Bromidum, Amyl Nitris, Analgen, Antipyrin, Antifebrin, Apomorph Hydrochloridum, Atropin Sulphas, Balsam Peruvianum, Bals Tolutanum, Belladonna, Caffeina Citras, Camphor, Cannabis Indic, Chloral, Chloroformum, Charta Nitrita et Chlorata, Cocaina Salicylas, Ethyl Nitris, Eucalypti Oleum, Euphorbia Pululiferi, Grindelia Robusta, Hyoscyamus, Iodipin, Lobelia, Myrrha, Nitroglycerin, Pilocarpina Nitras, Piscidia, Potass Bromid, Pot Iod, Potass Nitrat, Pyridin, Quebracho, Radium, Sodii Nitris, Spirit Ethera Nitrosi, Stramonium, Pulv Stramonii Comp, Tabaci Folia

Astringents —Section A

Bed Sores Acidum Boricum, Acid Sulphuric, Alum Sulph, Argenti Nitras, Amadou, Balsami Peruviani Ung, Brandy and white of egg mixture, Collodium, Plumbi Tannatis Glycerinum, Zinci Oxid Ung

Beri Beri Amyl Nitrite, Nitroglycerin, Saline Aperients, Strychnine

Bile, deficiency of Fel Bovinum, Hydrargyrum, Sodii Phosphas, Sodii Sulphas —See *Cholagogues* —Section A

Biliary Calculi —See *Gall stones*

Bites of fleas, to prevent —Lavand Ol, Pyrethri Flores (Insect Powder), Camphora

Bites and Stings of Insects (ants, bees, gnats, mosquitoes, wasps) —Chloroform, Ipecacuanha, Lotio Acid Carbolic, Liquor Ammonia, Liq Ammon Acet and Methylated Spirit, Liq Potassæ, Liq Sodæ, Liq Plumbi Subacetatis, Oleum Carbolisatum, Oleum Olivæ, Oleum Pulegi, Sodii Bicarb, all locally

Bites of Rabid Animals Acid Carbolic, Argent Nit, Caustery, Hydrag Perchlor
Bites of Snakes Acid Chromic, Caustery, Liquor Ammoniac, Potass Permang,
 Strych inject hypoderm, Tinct Ammon Comp, Anti-serum

Bladder, irritable Acidum Boricum, Belladonna, Buchu, Cannabis Ind, Chloral
 Hydras, Hyoscyamus, Opium *Mineral Waters* Fachingen, Malvern, Pougues,
 Lubatschowitz See also *Cystitis*, *Antilithus*, and *Urine*, in

Bleorrhagia — See *Gonorrhoea*

Blisters, to heal Unguent Cetacei

— to keep open Ung Mezelei, Ung Sabinae

Blood restorers — See *Tonics*, Section A

Boils *Internally* Acid Aisenios, Alkalis, Calx Sulphurata, Nuclein, Purgatives,
 Tonics, Yeast *Locally* Acid Carbolic, Glycerinum Belladonnae, Camphor
 Spu, Collodium, Galban Comp Ung, Menthol, Opium, Salol Camphor
Hypodermically Staphylococcus vaccine

Bones, Fracture of *Internally* Calcii Phosphas

Bowels, Torpidity of — See *Cathartics*

Brain, Inflammation of — See *Meningitis*

Breast, Inflammation of Glycerinum Belladonnae, *locally* Phytolacca, *inter-*
nally and locally

Breath, Fetor of Acid Carbolic, Carbo Ligni, Creosote, Oxygen, Potass Chloras,
 Potass Permang See also Antiseptic mouth washes, gargles and inhalations,
 Gastric tonics and intestinal antiseptics, laxatives, diphtheria, laryngitis, ozæna,
 phthisis, tonsillitis, etc

Bright's Disease, Acute Inflammatory Aconite, Ammon Acetat Liquor, Bella-
 donna, Calcii Chloridum, Cataplasmata, Digitalis, Diuretin, Elaterium, Tinct
 Ferri Acet, Jaborandi, Pulv Jalapæ Co, Juniper Ol, Leeches, Pilocarpin,
 Pot Acetas, Pot Tart Acid, Saline Purgatives, Scilla Scoparium, Spu
 Ætheris Nitrosi, Strontu Lactas

— *Chronic* Nitroglycerin Saline Aperients See also *Albuminuria*, *Dropsy*
(renal) and *Œræmia*

Bronchitis, Acute Acid Benzoic, Aconitum, Ætheri, Ammoniacum, Ammoniac
 Liquor, Ammonii Carbon, in large doses, Ammon Chloridum, Antim Tart,
 Apomorphinæ Hydrochloridum, Asafetida, Belladonna, Benzoin Ti Co,
 Tinct Camph Co, Chloral, Sp Chlorof, Cimicifuga, Copaiba, Croton Lin,
 Dionine, Eucalyptus, Ferri et Am Citras, Tinct Ferri Acet Ætheri, Galbanum,
 Heroin, Heroin Hydrochloride, Hyoscyamus, Iodipin, Ipecac, Larix, Lobelia,
 Potass Iod, Peronine, Plumbi Acet, Pulv Ipecac Co,
 Cataplasma, Strophanthus, Strychnina, Terebinth Ol,
 Yaba Santa

Bronchitis, Chronic Acid Benzoicum, Æthyl Iodidum, Ammoniaci Mist,
 Ammon Carbonas, Asafetida, Bals Peru and Tolu, Tinctura Benzoin Co,
 Caffein Citras, Chloral, Coninæ (Vapor), Codeinæ Syi, Creosoti Vapor, Cubeba,
 Oleum, Euphorb Pululif, Grindelia, Heroin,
 Iarg Pil, Hydrogeni Peroxidi Liquor, Hydrastis,
 Lobelia, Menthol, Morrhue
 Liq Syr, Pil Ipecac c Scilla,
 Pruni Ol, Pruni Sylvest Ol, Pot Iodid, Pruni Virginiana Syrup, Quillaria,
 Quinina, Scilla, Senega, Serpentina, Sulphur, Tar Water, Terebene, Tere-
 binthinæ Ol

Bronchocele or simple Parenchymatous Goitre Acidum Fluoricum Dil, Ammon.
 Fluorid, Hydrag Iodid Rub Ung, Iodoformum, Iodum, Potass Iodid,
 Sodn Iodidum, Thyroglandin, Thyroides Liquor, Thyroideum Siccum,

Brow Ague. — See *Neuralgia*.

Bruises Acetum, Acid Acetic Dil, Alum, Ammon Chlorid Lotio, Anthemis, Arnica, Calendulæ Flor, Calendulæ Tinctura, Capsicum, Hamamelis, Plumb Subacet Dil Liq, Saponis Linum, Soda Chlorid, Sp Vini Rect

Bubo, Acute Glycerin Belladonnæ, Iodoform, Lotio Acid Carbolici, Liquor Chlori and other antiseptic dressings

Burns Amadou Plaster, Cupri Oleatis Ung

Burns and Scalds Acid Borici Lotio and Ung, Acid Picric (Solutio), Acid Salicyl Lotio, Amylum, Benzoyl Peroxide, Oleum Carboheum, Bismuth Subnitrates, Calcis Lin, Calcu Carbonas Precip, Calcis Chlorin Liquor, Carron Oil, Cocaina, Colloidium, Ciccosotum, Creta Prepar, Eucalyptus Gauze or Oil, Flour, Gossypium, Iodoformum and Vaseline, Lini Oleum, Olive Oleum, Orthoform, Sp Rect *Internally* Stimulants, Digitalis, Morphine with caution

Bursitis, Acute Acid Carbol (inject), Blister, Tinct Iodi (paint or inject), Zinc Chlorid (inject)

Calculus, Ure or Lithic Acid, to counteract tendency to formation of—See *Antilithics*, Section A

—*Phosphatic*—See *Cystitis*, and *Urine, phosphatic*

Calculus, Renal—See *Urine, phosphatic, Antilithics*, Section A, and *Colic, Renal*

Cancer, Locally Acid Carbol, Acid Formic, Acid Nitric, Acid Sulph (Nordhausen), Glyc Acid Tannic, Antim Chloride, Arsenical Paste, Conium, Hydrag Nit Acid Liq, Iodoform, Methyl Violet, Pancreatic Enzymes, Potassa cum Calce, Potass Permanganas, Quinine Hydrochloride, Radium, Soda Cinamas, Soda Coumaras, Soda Meta coumaras, Soda Meta vanadas, Strontii Cinnamas, Violets, Zinc Chloridum *Internally* Acid Arsenios, Chelidonium, Chloral Hydras, Condurango, Exalgin, Methyl Violet, Opium, Orthoform, Pancreatic Enzymes, Terebinth Cina, Thalline Iodide, Thyroid Preparations, Uranium Salicylate

Carrunches—See *Boils*

Cardiac Tonics—See *Tonics*, Section A

Carminatives—Section A

Catarrh of the Respiratory Passages (common cold) Acid Carbolici (Vapor), Acid Salicylic (Vapor), Aconitum, Armoniacum, Ammon Benz, Ammon Chlor (Vapor), Sp Ammon Fetid, Amygdala Dulc, Antim Tart, Apomorphina, Bals Peruv, Bals Tolutanum, Benzoin Vapor and Insufflat, Bismuthi Subnitrat Insuf, Camphora, Cetraria, Cimicifuga, Cinchonidinæ Hydrobromidum, Dulcamara, Eucalyptus, Euphorbia Pululifera, Ferri's Snuff, Glycyrrhiza, Hordei Decoctum, Ipecacuanha, Linum, Lobelia, Menthol, Myrrha, Opium, Pinu Oleum, Pix Liquid, Pulv Ipecac Co, Quinine Sulph, Resorcin, Soda Chloridum, Smelling Salts, Senega, Sp Æther Nit, Syri Pruni Virg

—*Vesical*—See *Cystitis*

Cathartics—Section A

Caustics—Section A

Chafing of Skin Calamine Dusting Powder, Powdered Talc, Starch, Violet Powder

Chancres Acid Nitric, Acid Pyrogall, Acid Sulphuros, Argenti Nitras, Bismuthi Subiodid, Eucalyptol, Hydrag Lotio Nigra, Hyd Nitrat Liq Acid, Hyd Ox Rubr, Hydrogen Peroxid Liq, Iodoform, Iodol, Potass Permang, Resorcin *All locally*

Chapped Skin Amyli Glycerinum, Cerat Camphor, Glycerin Unguentum, Glycerin with Rose Water, Lanolin, Vaseline, Ung Aq Rose

Chilblains Acid Sulphuros, Alum Poultice, Amyli Glycerinum, Argenti Nitras, Aconit Lin, Belladon Lin, Boracis Ung, Bynes Ext, Calcu Chloridum, Calcu Lactas, Calcis Chlorinatæ Liq, Camphor Capsici Lin or Tinct Fort, Ciccosotum, Formaldehyde, Glycerinum, Ichthyol, Iodi Unguent, Moirhuæ Ol, Opn Lin, Saponis Lin, Ung Acid Carbolici, Ung Glyc Plumb Subacet, Toun

Cornea, Ulceration of Acid Boric, Argenti Nitras, Atropinæ Sulph. Liq., Atropin Ung., Hydrarg. Ox. Flav. Ung., Physostigmina Atropino to be used with caution in the elderly, for fear of glaucoma

Corns Acid Aceticum (Hæmule, Argenti Nitras, Collodium Salicylicum, Cupri Oleatis Unguentum, Plumbic Saponis Emp.

Corpulence—See *Obesity*

Coryza—See *Catarrh*

Cough Acid Hydrocyan. Dil., Acid Sulph. Dil., Acacia Gum, Agaricus, Amygdalæ Aqua and Mistura, Antim. Vinum, Apomorphina, Bals. Tolu., Benzoin Tinct. Composita, Codeina Sy. and Pastilles, Conium, Copaiba, Creosoti Vapor, Cubebs, Glycyrrhizum, Glycyrrhiza, Ipecacuanha, Lactuca, Linum, Lobelia, Menthol, Morphinæ et Ipecac. Troch., Opium, Piceidia, Pin. Liquidæ, Scilla, Styraç. Præp., Terebinthum—See also *Expectorants*, Section A

— *Spasmodic* Acid Hydrocyan. Dil., Belladonna, Cannabis Indica, Ammon. Bium, Tinct. Camph. Comp., Chloral Hydras, Cera. Oxalas, Conium, Hyoscyamus, Opium, Pruni Virg. Syrupus, Stramonium

Cramp—See *Antispasmodics*

Croup Anti diphtherial serum, Aconite, Alum, Antim. Tart., Apomorphina, Cupri Sulph., Emetics, Ipecacuanha, Lobelia. *Locally* Acid Lactic, Papain. *Externally* Camph. Linum Co., Cataplasmata

Cutaneous Diseases—See *Eczema, etc., etc.*

Cystitis Acid Benzoic, Acid Boric, Acid Camphoric, Ammonii Benzoas, Argenti Nitras (Injectio), Bellad. Supposit., Betol, Buchu, Cantharis, Capsicum, Cocaine Lactate, Collinsonia, Copaiba, Cystamine, Cubebs, Gmelina, Glusidum, Guaiacol Cinnamate, Helmitol, Hexamethylenetetramine, Hyoscyamus, Hydrag. Perchlor. (1 in 2000) irrigation, Kava Kava, Lysol, Methylen. Blue, Morphin. Suppos., Morphin. Ol., Naphthalene, Pænia, Potassii Benzoas, Potassii Citras, Potass. Licit., Pot. Permang. irrigation, Quinine, Salol, Santal. Flav. Ol., Sodii Benzoas, Sodii Phosphas Acid., Sodii Salicylas, Sulphaminol, Terebinth. Ol., Thymol, Tuticum, Tuberculin in tubercular variety, Uresin, Urotropine, Uva Ursi. *Laxatives*

Dandruff Borax Lotion, Hyd. Ammon. Ung., Oleum Carboicum, Paraffin Mollis, Sapo Mollis

Debility Acid Arsenios, Alcohol, Cajuputi Ol., Calumba, Chemical Food, Cinchona, Coca, Ferrum salts, Formates, Gentiana, Glycerophosphates, Hypophosphites, Lecithin, Morrhuæ Ol., Nuclein, Quassia, Quinine et Folia Citras, Strychnina. *Tonics*, Nervine and Stomachic

Delirium Antim. Tart., Belladonna, Cannabis Indica, Hyoscyaminæ Sulphas, Hyoscinæ Hydrobromidum, Methylal, Opium, Potass. Bromidum

— *Tremens* Ammoniac Liqueur, Amylen. Hydrate, Antim. Tart., Arnica, Cannabis Indica, Camphora, Camphora Monobrom., Capsicum, Chloral Hydras, Chloroformum, Digitalis, Hyoscyamin. Sulphas, Hyoscinæ Hydrobromidum, Hypnol, Normal Saline, Opium, Potass. Bromid., Scutellarin, Strychnina, Sulphonal

Demulcents—Section A

Depilatory Barri Sulphidum, Calx Sulphurata, X Rays

Desiccants—Section A

Diabetes Mellitus Acid Arsenios, Acid Lactic Dil., Acid Phosphor. Dil., Almond Cakes, Antipyrine, Arseni Bromidi Liqueur, Atropinæ Sulphas, Codeina, Creosotum, Eucalypt. Fol., Ferri Perchlor. Tinct., Ferri Phosphas, Glusidum, Guaiacol Benz., Hydrogeni Peroxidi Liqueur, Jambul, Lævuloso, Lithii Carbonas, Morphinæ, Opium, Pancreatin, Pilocarpina, Phosphorus, Potass. Citras, Potass. Permanganas, Sodii Bicarbonas, Sodii Phosphas, Sodii Salicylas, Strychnina, Suprarenal Gland, Urani Nitras. *Mineral Waters* Carlsbad, Vichy

— *Insuperdus* Ergota, Acid Gallic, Pot. Iod., Opium, Valerian

Dianthum, Acid Carboheum, Acid Gallicum, Acid Hydrochlor Dil, Acid Nitric Dil, Acid Phosph Dil, Acid Sulph Atom, Acid Sulph Dil, Acid Tannic, Alumen, Amylum, Argent Nitras, Bella Fructus, Bella Confectio, Bismal, Bismuthi Subnitras, Bismuthi Salicylas, Bismuthi et Cerni salicylas, Bismuthi Subgallas, Calori Carbon Præcip, Caloris Liqueor, Caloris Sacchi Liqueor, Cerebrum Essentia, Capsicum, Carbo Ligni, Catechu, Coton, Creosote, Creta Præp, Creta Atomat Pulv, P Creta Atomat c Opio, Cupri Sulph, Cholera Mixture, Doveri Pulv, Eucalypti Gummi, Ferrium salts, Fotonin, Glutanol, Granati Cort, Guaiacoli Valerianate Gumma, Hamamelis, Honthin, Hydragr Potchloi, Hydragr cum Creta, Kino, Linum, Naphthol, Opium, Plumbi c Opio Pil, Plumbi Aetas, Quinina Carbolis, Quinine Salicylas, Rosorin, Rhei Tinct, Ricini Oleum, Salol, Sassaparilla Medulla, Sinaruba, Dr Stevens' Pulvis Salinis, Tannigen

Diphtheria, Acid Carbolic Glycerin, Acid Formic, Acid Hydrochloric, Acid Lacticum (paint and spray), Acid Nucleic, and salts, Acid Sulphurous (spray) \ \ \ \ \ set \ \ \ \ \ or Liquor, Eucalyptol, Ferri \ \ \ \ \ Iodid, Iodoform, Iodine, Iodine, Iodine, Hydriodic Peroxide, Liquor, Iodi Tinct, Iodoform, Iodoform, Menthol Pigmentum, Papain (paint), Phenol Camphor, Potass Permanganas, Quinine Sulphas, Resorcin, Sode Chlorinate Liquor, Sulphur (insufflatio)

Diuretics —Section A

— *Hepatic* Ammon Chlor, Hydiarg Pil, Hydiarg Subchlor, Hydiarg Subchlor Co Pil, Juniperi Oleum, Taraxacum

Dysentery Acid Gallic, Acid Tannic, Alumen, Bellæ Confectio, Bismuth Cerium Salicylate, Cascariæ Infus, Catechu, Cubebæ Oleum, Cupri Sulph, Cuspariæ Infusum, Doveri Pulv, Gummi Rubrum, Hamamelis, Hæmatoxylinum, Hydrarg Perchlor Subchlor, Ipecacuanha, Lini Decoct, Naphthalene, Opium, Phenol Iodatum, Plumb Acet, Rhamni Sulphas, Rhamni Oleum, Salol, Salicylate, Samaruba, Sodæ Chlorin, Liq Soda Sulphas, T. T. T robinth Ol

Dysmenorrhœa Ammon Acetat Liquor, Amyl Nitris (inhalation),
 Apol, Belladonna, Boio-glyceride, Bromides, Cannabis Indica,
 Ergota, Guaiaci Resina, Plenacetin, Pulsatilla, Spirit Ætheris Nitrosi,
 Viburnum

Dyspepsia Acid Arsenios, Acid Carbohc, Acid Hydrochlor Dil, Acid Hydrocyan Dil, Acid Nitro-hydrochlor Dil Alcoh, Ammoniac Liquor, Ammonia Carbonat, Argem. Nitras, Bismuthi Caro Bismuti Subnitras, Buchu, Cera Caro Præparat, Calcei Liq, Calumna, Capsicum, Carbo Ligni, Caryophylli Ol, Cascar. P. Lin. Cerni Oxelas, Chiretta, Creosotum, Ferrum salt, Gentiana, Limonis Cortex, Magnesia, Magnesia Carbonas, Malt Extract,

Nux Vomica, Orexin Hydrochloride and Tannate, Papain, Pepsin, Peptonised Foods, Potassæ Liquor, Potass Bicarb, Potass Bichrom, Potass Sulph, Quassia, Quinina Sulph, Rheum, Salicinum, Sapo Durus, Senna, Serpentina, Sodæ Liq, Sodii Bicarb, Sodii Glycocholas, Sodii Sulphocarbolas, Sodæ Chlorin Liq, Somnifose, Strychnin Meta vanadas, Taka-Diastase, Taraxacum, Zingiber *Mineral Waters* Alet, Apollinaris, Chailottenbrunnen, Ems, Hom burg, Orezza, Vals — See also *Carminatives*, and *Tonics*, *Stomachic*, Section A

Dyspnœa — See *Asthma*, *Bronchitis*, *Cardiac Stimulants* (Section A), *Phthisis*, *Pneumonia*, *Rickets*, *Vaso dilators* (Section A)

Earache Almond Oil with Cocaine, Glycerinum, Morphine, or Opium Tincture

Ecbolics — Section A

Eczema Acid Arseniosum, Acid Carbolic, Acid Picric (Solutio), Acid Pyrogallie (oxidised), Acid Salicylic, Adops Linæ, Alkaline Solutions, Aluminium Oleas, Argenti Nitras, Aristol, Betulæ Oleum, Bismuthi Lotio, Cadmium Oleum, Calc Carbon Princip, Cunaphora, Chaulmoogia Oil, Creosoti Ung, Cremor Lithargyri, Cietæ Præp, Creolin, Dermatol, Episcarin, Euprophen, Gallanol, Glycerinum, Hyd Ammon Ung, Hydiarg Subchlor Ung, Ichthyol, Lassar's Paste, Lycopodium, Pepsinum, Picis Liquida Ung, Potass Carb (Lotio), Resorcin, Sodii Arsenas, Sodii Carb, Sodii Sulpho ichthyolum, Soroiodol, Tannoform, Ung Glycerin Plumbi Subacetatis, Zinci Oxidum *Mineral Water Aix les Bains*

— *Chronic* Acid Arseniosum, Ol Betulæ Ung, Cadmium Oleum, Hydiarg Nitrat Ung, Hyd Oxid Flav Ung, Naphthol, Paraffinum Liquid, Resorcin, Zinci Oxidum

Emetics — Section A

Emmenagogues — Section A

Emollients — Section A

Emphysema Ammon Carb, Digitalis, Iodipin, Nux Vomica, Pot Iodid, Pyridin, Quin Sulph, Saline Aperients

Epilepsy Acid Arseniosum, Æthylene Bromide, Ammon Bromid, Amyl Nitris, Amylene Hydrate, Argenti Nitras, Atropinæ Sulph, Auri Bromidum, Auri et Potassii Bromidum, Belladonna, Borax, Bromethylformine, Bromipin, Bromo hæmol, Camphora Monobrom, Castoreum, Cerii Oxalas, Chlorotone, Cupri Sulphas, Cypripedin, Ferri Perchlor Tinct, Ipecacuanha, Lithi Bromidum, Moschus, Niccoli Bromidum, Nitroglycerin, Opium, Picrotoxinum, Potassii Bromidum, Pot Iod, Rubidium Bromide, Rubidium Ammonium Bromide, Santonin, Sodii Bromid, Sodii Nitris, Spermin, Strontii Bromidum, Strychnina, Valeriana, Zinci Bromidum, Zinci Lactas, Z Oxid, Z Sulph, Z Valerianas

Epistaxis Acid Tannic, Adrenalin, Alum, Ergota, Galla, Gummi Rubri Extract Liquid, Ferri Chloroxydi Liquor, Hamamelis, Suprarenal Gland and Extract, Teiebinth Ol

Erysipelas Locally Acid Carbolicum (lotio), Acid Sulphurosum (spray), Amyli Glyce, Amylum, Argenti Nitras, Belladonnæ Glycerinum, Collodium, Creosotum, Guaiacolinol, Ichthyol, Iodi Liquor Fortis, or Ung, Lycopodium, Plumbi c Opio Lotio, Salol, Thiol *Internally* Aconitum, Belladonna, Cinchona, Ferri Perchlor Tinct, Guaiacol, Lactophenin, Quinina

Escharotics — Section A

Evacuations, Fetid Acid Carbolic, Bismuth Beta naphthol, B Salicylate, Calomel, Salol, Sodii Salicylas, Potass Permangan, Sodæ Chlorinatæ Liquor, Stomachic Tonics and Intestinal Antiseptics

Excoriations Alum, Acid Boric, Amylum, Boracis Glycerinum, Calamina Præp, Fuller's Earth, Glycerini Ung, Plumbi Carb, Zinci Oxid

Expectorants — Section A

Expectoration, Fetid Acid Carbolic, Chlori Liq, Creosotum Potass Permanganas, Menthol and Guaiacol (by intralaryngeal injection)

Fortis, Iodī Tinct (inject), Iodoform, Morrhue Oleum, Potass Iodid, Lin Potass Iodid c Saponē, Sodæ Chlorinatæ Liquor *Mineral Waters* Kœnigsdorff, Leuk, Marienbad

Gleet — See *Chronic Gonorrhœa*

Glycosuria — See *Diabetes Mellitus*

Gothæ, Simple Parenchymatous — See *Bronchocele*

— *Exophthalmic* Ammon Picras, Arsenical Liq, Digitalis, Ergot, Ferum, Iodī Tinct, Hyd Iod Rub Ung, Opium, Sodū Phosph, Strontū Bromidum, Strophanthus, Supraioral Gland, Thymus tablets Thyroid should never be given

Gonorrhœa, Acute Internally Aconitum, Antim Tut, Gonol, Hordei Decoct, Hyoscyamus, Lini Inf, Methylene Blue (Pure), Pareira, Potass Bicarb, Santal Flav Ol, Suintyl Locally Actol, Alumen, Argentamin, Argenti Nucleas, Argoun, Argynol, Borax, Betol, Bismuth Subnit, Cumin, Cupri Sulphocarbolas, Glibronol, Hydrag Nucleas, Ichthargan, Itrol, Iodoform and Eucalyptus Bougies, Largin, Novargan, Potass Permanganas, Protargol, Sodū Chloridum, Zinci Acetas, Z Chlorid, Z Permang, Z Sulphocarbolas

— *Chronic, or Gleet Internally* Arhovin, Copaiba Cubeba, Diptercarpī Balsamum, Ferri Perchlor Liq freely, Gonol, Magnes Sulph, Santalū Oleum, Triti cum Locally Acid Tannic, Argenti Nitras (bougie), Cupri Sulphas, Dextroform, Plumbi Acetis cum Opio, Quercus, Zinci Acetas, Z Chloridum cum Belladonna, Z Sulphas

Gout Acid Aconiosum, Acid Chmic, Ammonii Chloridum, Ammonii Phosphas, Caffeina Di iodo-Hydriodidum, Capuputi Ol, Chaulmoogra Oil, Chinoline Periodide, Citum, Colchicum, Colchiemæ Salicylas, Colchi Sal, Crotonis Lin, Euonymin, Guaiaci Resina, Hydrag Pil, Hydrag Subchlor, Hyoscyamus, Lithium salts (see p 739), Lysidine, Lycetol, Magnesia, Magnes Sulph, Meotan, Morphinū Inject Hypod, Phenazonum, Piperazine, Piperazine Quinate (sidual), Piperidine Tartrate, Podophyllin, Potass Acetas, P Citras, Sabina, Salignum, Serpentina, Sodū Bicarb S Phosphas, S Taurocholas, Sodū Stlphas, Sparteine Periodide, Strontū Salicylas, Sulphur, Trimethylaminæ Hydrochloridum, Uresin, Uricedin, Urosine, Uystamine *Mineral Waters* Adelhheidsquelle, Aix les Bains Baden-Baden, Buxton, Carlsbad, Eilsen, Ems, Franzensbad, Ischia, Marienbad, Nenndorf, Neuenahr, Ofen, Plombières, Soden, Stathpeffer, Tarasp, Toeplitz, Vichy, Weilbach, Wiesbaden, Wildbad

Gout, painful Aconitine Unguent, Antipyrine, Capuputi Oleum, Hyoscyamus, Menthol, Morphina, Opium, Potass Iodidum, Veratrinæ Unguentum

Gums, inflamed Alumen, Boracis Glycerin, Gummi Rubri Tinct, Krameria Tinct, Myrrha Tinct, Myrrha et Boracis Tinct, Potassū Chloras, Pyrethri Tinct, Quercus Decoct

Hæmatemesis Acid Gallicum, Acid Tannicum, Alumen, Argent Nitras, Ergota, Ferrie salts, Hamamelis, Ico, Morphin Inj Hypod, Opium, Plumbi Acetas, Sodū Chloridum, Terebinthinæ Oleum

Hæmatinics — Section A

Hæmaturia Acid Sulph Dil, Alumen, Ergota, Ferri Perchloridi Liquor, Hamamelis, Plumbi Acet, Terebinthinæ Oleum

Hæmophelia Adrenalin, Calcū Chlorid and Lactas, Ergot, Hamamelis, Terebinth Ol

Hæmoptysis Aconitum, Amyl Nitris, Calomel, Calcū Chloridum, Digitalis, Eucalyptol, Ferri Acetatis Liquor, Gelatinum, Hamamelis, Hydrag c Creta, Ipecac, Ipecac c Opio P, Sodū Chlorid, Morphina, Opium, Plumbi c Opio Pilula, Saline Purgatives, Terebene, Terebinth Oleum, Terpene Hydrate

Hæmorrhage — See *Styptics*

— *Post partum* Copious intra uterine irrigation with water at 118° F, Ergota, Ergotine (Inj Hypod) Ergotinine, Normal Saline Solution (Transfusion)

— *Uterine* — See *Menorrhagia*

Hæmorrhoids Acid Nitricum (lotio), Acid Tannic Ung, Aesculet Anusol, Belladon Ung, Calomel, Cascara Sagrada, Cetaceæ sine Benzoin Ung, Coni Ung, Galban Ung Co, Gallæ Ung and Ung cum Opio, Glycyrr Pulv Co, Iod. Form (Supp), Morphina, Picis Pilulæ et Capsulæ, Piper Nigrum, Sennæ Confect, Stramon Ung, Sulphur, Mineral Waters, Mergentheim.

Hæmostatics —Section A

Hair falling off See Alopecia

Hay fever Acid Chromic, Adrenalin, Belladonna, Camphor, Carbon Tetrachloride, Eucalypti Oleum, Grindelia, Menthol, Potass Iodid, Quinina Sulphas Acidus, Stramonium, Carbolic Smelling Salts, Suprarenal Gland and Extract, Zinc Phosphidum, Zinc Valerianas

Headache, Internally Acetanilide, Acid Hydrobrom Dil, Ammon Bromid, Ammon Aromat Spirit, Amyl Nitris (vapor), Antipyrine, Cannabis Ind, Caffeina, Camicifuga, Lactophenium, Magnesia, Nitroglycerin, Phenacetin, Potass Bromid, Pot Iod, Quinina Sulphas, Sodii Bicarb, Sodii Salicyl Locally Aconitum, Æther, Cocaine, Menthol, Opi Lin

Heart, Valvular Disease of Adonis Vernalis, Apocynum, Æthoxycaffeinum, Caffeina, Convallaria, Digitalis, Erythrophlæum, Sparteina, Strophanthus

Heartburn —See Pyrosis

Hectic Sweating —See Sweating

Hepatics —See Cholagogues Section A

Hepatitis Acid Nitro-hyd Dil, Ammon Carb, Hydrarg, Hyd Iod Rub Ung Ipecacuanha, Potass Iodid

Herpes Internally Morphina Tart (hypod inj), Potass Iodid, Purgatives, Locally Acid Boric, Amyl Glycerinum, Argenti Nitias, Ammon, Menthol, Zinc Ung, Unna's Zinc Gelatin

Hiccough Ætheris Spt, Amyl Nitris, Blister over Cervical, Bromides, Camphor, Chloral, Chloroform Spt, Ergota, Morphina, Nitroglycerin, Pilocarpina, Sinapis Infusum, Terebinth Ol 3i, Zinc Valerianas

Hydrocele Acid Carbolic, Glycerinum and Tinctura Iodi

Hydrocephalus Crotonis Oleum, Hydrarg, Subchloridum, Potass Bromidum, Potass Iodidum

Hyperæmia Cannabis Indica, Chloral Hydras, Chloroformum, Curara, Morphina

Hypnotics —Section A

Hypochondria Acid Nitro-hydrochlor Dil, Cholagogues and Chloral Hydras, Nerve Tonics, Potass Bromidum, Strychnina Mineral Water Homburg

Hysteria Ammonia Fetidus Spiritus, Ammon Carb, Ammon Bromid, Ammon Valerianas, Asafetida, Auri Bromidum, Auri et Potass Bromid, Camphori Ol, Camphora Monobromata, Tinct Chloroformi et Morphina Co, Ferrum salts, Lavand Ol, Valerianate, Moschus, Nux Vomica, Phosphorus, Potass Bromid, Quinina Sulph, Rosmarini Ol, Ruta Ol, Strychnina, Sumbul, Terebinthina Ol, Valeriana, Zinc Phosphidum, Zinc Valerianas Mineral Waters Homburg Lippik, Spa

Impetigo Contagiosa Hydrarg Ammon Ung, Iodoform Ung, Zinc Oleat Ung, Zinc Unguentum, all after removal of crusts by soaking in oil, or starch poulticing

Incontinence of Urine —See Urine.

Indigestion —See Dyspepsia.

Inflammation Acute Aconite, Antim Tart, Belladonna, Glycer Bellad, Hydiarg Subchloridum, Opium

— *Chronic* Iodine and Iodides, Iothion

Influenza Acid Carbolic, Acid Sulphurosum (vapor), Ammon Acetat Liq, Antim Tart, Antipyrine, Benzoin Vapor, Benzol, Calx Sulphurata, Eucalypti Oleum, Eupatorium, Euquimine, Ipecac Co Pulvis, Phenocoll Hydrochloride, Potass Bicarb, Quinine Sulphas, Resorcin, Salicinum, Salipyrin, Sodii Salicylas, Sp Æther Niti, Tinct Quinine Ammoniatæ

Insects, to keep away Camphora, Colocynth Pulpa, Lavand Oleum, Menth Pip, Oleum Pyrethri Flores, Quassia, Rosmarini Oleum, Terebinth Oleum

Insomnia —See *Hypnotics*, Section A

Iritis Acid Boric Lotio (hot), Atropine Guttæ or Ung, Atropin Methyl bromid, Belladonna, Canthar Emp, Cocaine, Duboisine, Hydrarg Perchlor and Subchlor, Hyalidine, Hyoscine, Pilocarpin Nit Inj Hyp, Potass Iodidum, Pulv Doveri, Quinine

Irritants —Section A

Itch —See *Scabies*

Itching —See *Pruritus*

Jaundice Acid Nitro hydrochlor Dil, Alkalis, Aloes, Ammonii Chlorid, Creosotum, Euonymum, Fel Bovinum, Hydiarg Subchlorid, Iodin, Pilocarpina, Potassa Sulphurata, Podophyllin, Potassu Sulphas, Sapo Durus, Sodii Sulphas, Taraxacum

<i>Joints, Rheumatic</i>	}	Belladonna Emp, Hydrarg Oleas, also with Morphia, Ung
— <i>Enlarged</i>		Hydrarg Comp, Iodum, Lin Potass Iod c Saponæ, Potass Iodid, Plumbi Iodidi Ung, Salocrocol, Sodii Salicylas, Veratrine Ung

Kidney Disease —See *Albuminuria*, *Bright's Disease*, *Dropsy (Renal)*, and *Œræmia Contra indicated* Opium, Cantharides, Turpentine

Laryngismus Stridulus Amyl Nitris, Antipyrine, Belladonna, Chloral Hydras, Chloroformum, Potassu Bromidum, Ricini Ol, Rheum, Hot Water

Laryngitis Aconiti Tinct, Antim Tart, Codeina, Guaiacum *Locally* Acid Lactic, Acid Sulphuros (spray), Acid Tannic Glycerin, Alum, Ammonium Chloride, Argenti Nit, Belladon Glyc, Benzoin Vapor, Closooti Vapor, Ice, Menthol (spray), Pini Sylvest Oleum

Laxatives —Section A

Leech bites, to stop bleeding from Alum, Argenti Nitias, Collodium, Ferru Perchlor, Matico, Ol Terebinth

Leeches, to dislodge if swallowed Sodii Chloridum, in strong solution

Leprosy Balsam Dipterocarpi, Chaulmoogra Oil

Leucocythæmia Acid Arseniosum, Bone Marrow, Ferrum salts, Lecithin, Phosphorus

Leucorrhœa Acid Boric, Acid Carbolic, Acid Chromic, Acid Gallic, Acid Tannic, Alumen, Bismuth Subnit, Borax, Cantharis, Catechu, Creolin, Cupri Sulphas, Cyllin, Ferrum salts, Granati Cort, Gummi Eucalyptus, Hamatoxyl Decoct, Hydrarg Perchlor, Krameria, Pareira, Potass Iodidum, Quercus Cort, Quinine Hydrochlor, Santal Flav Oleum, Sodii Sulphocarbolas, Tomes, Zinci Sulph, Zinci Sulphocarbolas *Mineral Waters* Kreuznach, Wildungen

Lice —See *Pediculosis*

Lichen Planus Locally Acid Carbolic, Acid Hydrocyan Dil, Acid Sulphurosum, Hydiarg Oxid Flav Ung, Ichthyol, Pix Liquida, Zinci Ung *Internally* Antimony, Arsenic, Hydrargyrum

Lips, cracked Adeps Lanæ, Bals Peru Unguent, Cetacei Ung, Paraffinum Molle

Mumps Aconitum, Belladonna Glyc, Doveri Pulv, Hydrarg cum Creta, Jaborandi, Opium, Pilocarpina

Myxædema Thyroidei Liquor, Thyroideum Siccum, Thyroglandin

Nerv Acid Chromic, Acid Nitric, Alum, Liq Ferri Perchlor Fort, Liq Sodii Ethylatis, Zinc Chloridum, Zinci Nitratis

Narcotics —Section A

Nausea —See *Vomiting*

Nephritis —See *Kidney Disease*

Neuralgia Acetanilide, Acid Arsenios, Acid Osmic, Aconiti Chloroform, Aconiti Linim, Aconitinæ Ung, Aconitum, Æther (spray), Æthoxycafeinum, Ammon Bromidum, Ammon Chlorid, Ammon Valerianas, Amyl Nitris, Amygdophenim, Analgin, Antikamnia, Antitoxine, Atropinæ Solut (hypodermically), Atropinæ Valerianas, Belladonnæ Lin, Butyl Chloral Hydras, Caffena, Camphore Lin, Camphor Lin Ammon, Cannabis Indica, Canthar Emp, Carbon Tetrachloride, Chloral cum Camphora, Chloroformum, Cimicifuga, Cinchona, Cocaina, Conium, Crotonis Liniment, Delphinina, Eucalgn, Ferri salts, Gelsemi Tinctura, Gelsemum, Guaiacol, Hyoscyamus, Iodoform, Kryofin, Lactophenim, Malakin, Menth Pip Oleum, Menthol, Methyl Chloridum, Mesotan, Migranine, Morphina, Morrhue Ol, Opium, Papaveris (Dococtum), Phenacetin, Phenazonum, Phosphorus, Piscidia, Quininæ Glycerophos, Quininæ Sulph, Salophen, Salicin, Scutellarin, Sinapis (Cataplasma), Sodii Salicyl, Strychnina, Veratrinæ Ung, Zinci Valerianas

Neurasthenia —See *Debility*

Nipples, Sore or Fissured Acid Sulphurosum, Acid Tannic Glycerinum, Argent Nitras, Bals Peru Ung, Boiacis Ung, Catechu, Orthoform, Plumbi Tannatis Glycerinum, Sodii Chlorinat Liq

Nitrates of Silver stains, to remove Potass Cyanid, Potass Iodid, Sodium Thiosulphate

Nocturnal Emissions Belladonna, Ferri Bromid, Potass Bromid

Nymphomania Ammon Bromidum, Camphora, Chloral, Conium, Potass Bromidum

Nutritives —Section A

Obesity Alkalis, Ferri Iodid, Fucus Vesiculosus, Potass Iodid, Thyroideum Siccum *Mineral Waters* Carlsbad, Ems, Kissingen, Marienbad, Tarasp

Ophthalmia Neonatorum Acid Boric, Alum, Argentamin, Argent Iodid, Argent Nit, Argyrol, Cocaine, Collargol, Cuprargol, Cuprol, Hydrarg Cyanid, Hydrag Ox Flav Ung, Hydrag Perchlorid, Iodoformi Ung, Liq Calcis Chlorinat, Mitigated Caustic, Protargol, Quinin Sulphas, Tachiol, Zinci Sulphas

Orchitis, Acute Locally Glycerinum Belladonnæ, Plumbi Acet et Opi Lotio *Internally* Antimonium Taiturat, Guaiacol, Hyoscyamus, Phenylurethane, Phytolacca, Saline Aperients

Otorrhœa Acid Borici Lotio, Iodoform, Iodol, Potass Permang, Zinci Chlorid *all locally*

Ozena Acid Carbohic, Acid Chromic, Acid Boric, Borax, Boro-glyceride, Creosotum, Iodoform, Menthol, Potass Permanganas, Sodii Chloridum, Sodæ Chlorinat Liquor, Sodii Ethylatis Liquor, Thymol, Zinci Chlorid, *all locally*

Palpitation Acid Arsenios, Acid Hydriobrom, Acid Nitro hydrochlor, Aconitum, Æther, Ammonia, Belladonna, Bromides, Anti dyspeptic remedies, Ferrum salts, Hydrarg Pil, Syr Acid Hydriod, Strychnina

Paralysis (Peripheral and Functional) Belladonna, Cannabis Ind, Ergota, Ferrum salts, Hyoscyamus, Nux Vomica, Phycostigma, Strychnina *Mineral Waters* Aix-la Chapelle, Baden Baden, Eilsen, Ischia, Kieumach, Toeplitz

— of *Lead Poisoning* Alkaline Sulphates, Potassum Iodidum

Parasites, Intestinal — See *Anthelmintics*, Section A

Pediculosis Acetum (warm), Bals Peruvianum, Hyd Ammon Ung, Hydrag Oleas, Naphthol, Oleum Carbolicum, Paraffin Oil, Resorcin Camphor, Sas-afras Oil, Staphisagriae Olei Ung, Styriacis Ung, Sulphur Ung

Periostitis Counter-irritants, Iothion, Potassu Iodidum, Vesicants

Peritonitis, Acute Belladonna, Hydrag Subchlor, Opium, Iodine

Perspiration, to diminish — See *Anhidrotics*, Section A

Perspiration, Fetid Acid Boric, Acid Carbolie, Acid Salicylic Glycer, Belladonna, Plumbi Oxid Ung, Pulv Salicylic cum Talco, Salicylic Subst, Zinci Ung

Phthiasis — See *Pediculosis*

Phthisis Acetophenone (inhal), Acid Benzoic (inhal), Acid Carbolieum, Acid Cinnamic, Acid Hydrocyan Dil (inhal), and salts, Acid Tannicum, Acid Hydrofluoric (inhal), Aconiti Tinct, Agaricin, Alcohol Methyleum, Antifebrum, Aniline, Atropina, Camph Tinct Co, Carbon Bisulphidum, Cicuton, Creosotum, Creosote Carb, Guaiacum, Guaiacum Ole (inhal), Guaiacum Chloroform, Formaldehyde, Guaiacum, G Benzoate, G Camphoric, G Carbonate, G Cinnamate, and other Guaiacol salts, Guaiacyl, Guaiacum, Helenin, Heroin, Heroin Hydrochloride, Igazol, Iodi Vapor, Iodoform, Lachnanthes, Malti Extractum, Menthol and Menthosol, (intralaryngeally), Morrhua Oleum, Opium, Pancreatic Emulsion, Peronine, Piperidine Guaiacolate, Pneumin, Pilocarpinae Phenas, Pini Oleum (vapor), Plumbi Acetas, Pruni Virgin Syr, Quinina and Quinine Salts, Radium, Saccharum Lactis, Salol, Sodii Cacodylate, Sodii Metaphosphates, Sodii Cinnan, Sodii Hypophosph, Sodii Meta-vanadas, Sodii Thiochol Tinctulin, Urea

Piles — See *Hæmorrhoids*

Pleuritis Acid Aceticum, Acid Boric, Acid Salicyl, Argent Nitrosi Sp, Glycerinum, Cadinum Oleum, Hydrag Oleas, Hydrag Oxid Rub Ung, Naphthol, Piceis Ung, Resorcin, Sodii Hyposulphur, Zinci Ung

Plague Acid Carbolie, Anti-plague serum, Glyc Belladon, Calomel, Stimulants, Strychnina

Pleuritis Aconitum, Antum Tart, Canthar Emp, Cicutonis Linim, Hydrag Potass Iod, Sinapis Cataplasma, Diaphoretics,

Pneumonia Aconitum, Aconitum Acetat Liquor, Ammon Carbonas, Amyl Nitrosi, Ant Nitrosi Sp, Caffeina, Calci Chloridum, Canthar Empl, Carbonis Bisulphid, Digitalis, Helenin, Heroin, Heroin Hydrochloride, Iodoform, Moschus, Oxygen, Potass Bicarb, Quinina, Sinapis Cataplasma, Strophanthus, Strychnine, Diuretics, Diaphoretics Cathartics

Polypi, Nasal Locally Acid Chromic, Acid Tannic, Absolute Alcohol, Sodii Ethylatis Liquor, Zinci Chloridum

Post partum Hæmorrhage — See *Hæmorrhage, Post partum*

Prolapsus Ani Acid Tannic, Alum, Capri Sulph, Ligotin, Ferri Perchlor, Gummi Rubr Extr Liq, Hamamelis, Nux Vomica, Quercus, Sulphur

Prostration Aether, Ammonia, Caffeina, Coca, Moschus, Nerve Tonic, Spiritus Vini Gallici Mistura, Strychnina

Pruritus or Itching Internally Acid Arsenos, Ammon Bromid, Hyoscyamus, Quinina Sulfurina *Locally* Acid Boric, Acid Carbolie, Acid Hydrocyan Dil, Argenti Nit Bromid Subst, Borax, Cocaine, Urea Gallica, Cupri Sulphas, Glycerin Ichthiol Iodoformum, Papaveris (Decoctum), Pectus Bals, Lauracerasi Aq, Liquor Carbonis Detergens, Plumbi Subacet Liq, Sodii Bicarb Lotio, Sulphuris Ung, Zinci Ung

Pruritus Ani Acid Carbohc Ung, Acid Salicylic Ung, Gallæ c Opio Ung, Hydrarg Subchlor Ung, Menthol, Pix Liq, Plumbi Acet, Purgatives, Resorcin

— *Vulvæ* Aluminium Nitrate, Glycerinum Boracis, Cocaina, Ichthyol, Pilocarpine Nitrate, Carbonis Deterg Liq, Menthol, Plumbi Subacet Liq

Psoriasis Internally Acid Arsenios, Atoxyl, Dulcamara, Salicin, Thyroideum Siccum *Locally* Acid Carbohc, Acid Pyrogallie Oxydat, A Salicylic, Anthrubicu, Aristol, Betulæ Albæ Olei Ung, Chaulmoogra Oil, Chrysarobinum, Creosotum, Dulcamara (Decoct), Epican, Gallanol, Glycerinum, Hydrarg Sozonodolas, Hydring Subchlor, Hydracetin, Ichthyol and Compounds, Liquor Carbonis Detergens, Naphthol, Ol Cadinum, Picis Unguent, Potassa Sulphurata, Potass Iodidum, Radium, Resorcin, Saponis Emp, Sodii Carbonas

Puerperal Convulsions Chloral, Chloroformum (inhal), Morphina, Normal Saline, Potassii Bromidum, Thyroid prep

Purgatives — Section A

Purpura Arsenic, Ferri Perchlor Tinct, Hyd c Creta, Hyd Perchlor, Calciu Chlorid, Quinina, Sodii Salicylas, Terebinthine Ol or other intestinal antiseptics

Putrescence, to Correct — See *Antiseptics*, Section A

Pyæmia Alcohol, Ammonia, Antiseptics, Antistreptococcus or Antistaphylococcus serum, Quinina

Pyrosis Acid Hydrochlor Dil, Acid Sulphuros, Argent Oxid, Bismuthi Subnitras, Bismuth Carb, Catechu, Ceru Oxalas, Magnesia, Manganesi Oxid Prep, Opium, Pulvis Doveri, Sodii Bicarb, Sodii Sulphocarbolas

Refrigerants — Section A

Restoratives — Section A

Rheumatism Acute Acid Salicylic, Acid Benzoic, Aconitum, Acetanilide, Acetopyrine, Amygdophen, Antirheumatic, Betol, Canthar Emp, Cimicifuga, Gaultheria Ol, Limonis Succus, Opium, Methyl Acetyl Salicylate, Methyl Salicylate, Mesotan, Phenazone, Phenocoll Hydrochloridum, Pot Acetas, Pot Bicarb, Potass Citras, Pulv Doveri, Pipelazine Quinate, Pyramidon Salicylate, Quinina, Rheumatine, Salicinum, Saligenin, Salt, Salocoll, Salol, Salophen, Sodii Diethosalicylas, Sodii Salicylas, Tolypyrin, Tolsal, Trimethylaminæ Hydrochloridum

— *Chronic* Acid Acetylsalicylic, Acid Arseniosum, Acid Formic, Acid Salicylic, Aconiti Lin, Aletris, Ammon Chlorid, Ammon Phosp, Antum Sulphurat, Armoracia, Asaprol, Bellidonn Lin Co, Betol, Buchu, Camphor Ol Essent, Capsici Tinct Fort, Chelsea Pensioner, Chloral, Chloroformum Camphoratum (local), Conium, Cajuputi Ol, Chaulmoogra Oil, Citarin, Citrophen, Crotonis Oleum, Dulcamara, Fluorrhæum, Guaricol and G Carbonas, Guaiacum, Hydrarg Iodid Rub, Hydrarg et Morphina Oleas, Iodi Liquor Fortis, Iodipin, Ichthyol, Iodoform, Limonis Succus, Lin Camph Co, Lithii Guaiacas, Lithii Salicylas, Lycetol, Lydine, Magnesia, Malakin, Menthol, Mesotan, Methyl Acetyl Salicylate, Methyl Salicylate, Morrhuæ Oleum, Myristicæ Oleum, Opium, Phenacetin, Phenazonum, Picis Burgundicæ Emplast, Pini Oleum, Pini Sylves Ol, Piperazine Quinate, Potassa Sulphurata, Potass Iodid, Lin Pot Iod c Sapon, Lin Saponis, Pyramidon Salicylate, Syr Quinina Hydrobromidum and Hydriodidum, Salpyrin, Salol, Sarsaparilla, Serpentina, Sodii Iodid, Sodii Salicylas, Strontii Salicylas, Sulphur, Terebinth Lin, Trimethylaminæ Hydrochloridum *Mineral Waters* Aix les Bains, Aix la Chapelle, Barèges, Baden Baden, Bath, Berka, Buxton, Franzensbad, Lucca, Ofen, Toeplitz, Wiesbaden, Woodhall

— *Painful* Belladonnæ Chloroformum, Hydrarg et Morphina Oleas, Lin Camph Comp

Rickets Acid Phosphor Dil, Calci Liquor, Calciu Chloridum, Calciu Phosphas, Creta Preparata, Ferri Phosphas Morrhuæ Oleum, Chemical Food, Thyroid preparations,

Spermatorrhœa Belladonna, Camphor, Camphor Monobromata, Capsicum, Ferrum salts, Nux Vomica, Potassum Iodidum, Purgatives, Quinina, Stychnina

Syphilis Bifida Iodo Glycerin Solution (Morton's) injected

Sprains Aconit Lin, Bellid Lin and Emp, Calendula, Carbolic Fomentation, Cold Douche, Iodi Liq Fort, Opii Linim, Lin Saponis, Sp Vini Rectif (lotion), Terebinth Lin Acet, Sodii Chloridum (Fomentation)

Stimulants—Section A

Stings—See *Bites and Stings*

Stomach Pain—See *Gastralgia and Tonics*, *Stomachicæ*, Section A

—*Ulceration of* Bismuth Carb, Ferru Sulph, Ico, Magnes Carb, Magnes Sulph, Morphin Inj Hypod, Opium, Peptonised Foods, Potass Bicarbonas, Sodii Bicarb

Stomachics—Section A

Stomatitis, Ulcerative Alum, Borax, Boracis Glycerin, Tinct Myrrhæ et Boracis, Potass Chlorat

Strangury Belladonn Suppos, Camphor, Morphin Hypod Inj and Suppos

Styptics—Section A

Sudorifics—Section A

Sunstroke Apomorphin, Atropina, Cold Douche, Phenazonum, Purgatives, Sinapisms

Sweating, to diminish—See *Anhidrotics*, Section A, and *Fetid Perspiration*, Section B

Syncope Æther, Ammon Spir Arom, Sputus Vini Gallici

Synovitis Blister, Emp Ammon c Hydrarg, Hydrarg Oleas, Ung Iodi, Tinct Iodi (inject), Pigment Iodi Co

Syphilis Primary and Early Secondary Acid Chromic Pigmentum, Auri Chloridum, Barium Chloride, Calomel Cream, Hydrargyrum and its Compounds, Mercurial Cream, Rubidium Iodide, Stillingia and Fluid Extract

—*Late Secondary and Tertiary* Auri et Sodii Chloridum, Iodipin, Iodum and the Iodides, Hydrag Carbolas, Hydrag Soroiodolas, Morrhua Oleum, Quinine Iodo hydriodide, Sajodin, Sarsaparilla *Mineral Waters* Aix-la Chapelle, Kreuznach, Vals, Woodhall

Syphilitic Nodes Emplastum Hydrargyri, Potassum Iodidum, Sodii Iodidum

—*Warts*—See *Warts*, *Syphilitic*

—*Ulcers* Ung Amyli Iodidi, Hydrarg Nit Liquor Acidus, Iodoformum, Causticum Iodi, Hydrarg Flava or Nigra Lotio, Hydrag Perchlor Lotio

Tabs Mesenterica Hydrarg Liniment and Oleas, Iodoformum, Morrhua Oleum, Ferrum preparations of, Quinina

Tape Worm—See *Anthelmintics*, Section A

Teeth, Caries of Acid Carbolic, Arsenical Paste, Cocaine, Chloral cum Camphora et Cocaine, Creosotum, Mastic Dentaire

Tetanus Acid Carbolic, Amyl Nitris, Atropina, Cannabis Indica, Chloral Hydras, Curara, Magnes Sulphas, Physostigma

Thirst, to allay Acid Citricum, Acid Phosphoricum Dil, Acid Sulph Aromat Acid Tartaricum, Imperial Drink, Limonis Succus

Throat, Sore—See *Tonsillitis*

Thrush—See *Aphthæ*

Tic Douloureux—See *Neuralgia*

Tinea Capitis—See *Ringworm*

Tonics—Section A

Tonsils, Enlarged Internally Potass Iodidum *Locally* Acid Carbolic Glycerin, Acid Tannic Glycerin, Ferr Perchlor Glycel, Iodum cum Glycerino or Tinct Iodi

— *After Excision of* Trochiscus Althææ

Tonsillitis Internally Aconitum, Antim Tart, Hydrarg c Creta, Ipecac Pulv Co, Sodii Salicylas, Purgatives *Locally* Acid Acetic, Acid Carbolic Glyc, Acid Sulphuros, Acid Hydrochlor Dil, Acid Tannic Glyc, Alum, Argenti Nitras, Boracis Glyc, Capsicum, Catechu Troch, Cocaina, Chlori Liquor, Cubebe Troch, Gummi Eucalypt Troch, Ferr Perchlor Ti and Glycerin, Hydrarg Perchlorid, Kîmmerl, Myrrha, ~ Mandl, Pot Chloras, Pot Nitras, Pot Permang, Rosæ Inf Acid, Sodæ Chlorinatæ Liq

Toothache Acid Carbolic, Acid Sulphuros (spray), Aconite and Iodine, Cajuputi Oleum, Capsici Tinct Fortior, Caryophylli Oleum, Chloral cum Camphora et Cocaina, Chloroform c Camphora, Croosotum, Eucalgin, Gelsemii Tinctura, Menthol, Phenol Camphici, P'p'p'p', Quinine Ammoniatæ Tinctura

Trichiniasis Glycerin in large doses, Torobinth Ol, Purgatives

Tuberculosis Barium Chloride, Calci Chloridum, Calci Lactas, Calci Phosphas, Calx Sulphurata, Ferr Iodid, Ferr Phosphatis Syrup Co (Chemical Food), Galium Sparagacis, Hyd Subchlor, Iodum, Moribundæ Oleum, Potass Iod, Potass e Quinæ Sulph, Sodii Iodidum *Mineral Waters* Barèges, Caunterets, Ems, Ischia, Koenigsdoiff, Kosen, Krankenhêil, Kreuznach, Luhatschowitz, Neuenahr, Reichenhall, St Moritz, Soden, Strathpeffer, Vals, Woodhall — See also *Phthisis*

Typhoid Fever — See *Fever, Typhoid*

Ulcers, Heal ~ ~ ~ Acid Boric, Argenti Nitras, Bals Peruv, Bismuth ~ ~ ~ Carbon Piæcip, Calci Iodidum, Calcis Chlorinatæ ~ ~ ~ Creta Præparat, Cupri Sulphas, Iodipin, Orthoform, Plumbi Acetas, Plumbi Carb, Resinæ Emp, Sabina, Zinci Sulphas, Zinci Ung

— *Malignant* Acid Chromic, Acid Nitric, Antim Chlor Liquor, Potassa Caustica, Radium, Zinci Chlorid

— *Foul* Acid Carbolic, Acid Chromic, Acid Lactic, Acid Salicylic, Acid Sulphuros, Argenti Nitras, Bismuthi Subiod, Calcis Chlorinatæ Liquor, Carbo Ligni, Chlori Liq, Calx Chlorin, Cinchona, Cupri Subacetat, Eucalypti Ung, Hydrarg Perchlor Lotio, Iodoform, Potassa Caustica, Potass Permanganas, Resorcin, Sodæ Chlorinatæ Liquor, Zinci Chloridum

— *Indolent or Callous or Chronic* Acid Chromic, Alumen Exsic, Argent Nit, Bals Peruvianum, Benzoini Tinct Co, Cupri Acetas, C Subacetat, C Sulphas, Elemi Ung, Hydrarg Lin, Hydrarg Oxid Rubr Ung, Ichthyol, Kino, Lotio Rubra, Pepsinum, Sabina Ung, Unna's Paste, Zinci Chloridum

Uremia Aconite, Amyl Nitris, Caffeina, Digitalis, Pulv Elateri Co, Pulv Jalapæ Co, Jaborandi, Magnes Sulph, Nitroglycerin, Normal Saline Solution (12 c f - 100) ~ ~ ~ ypodermically, Potass Acet, Potass Bicarb, Spt Chloroform for the convulsions

Urethritis Alkalis, Saline Purgatives, Oleo-Balsams, Alcohol interdicted — See also *Gonorrhœa, acute and chronic*

Urine, Alkalinity of — See *Urine, Phosphatic, and Cystitis*

— *Deposit of Uric Acid or Urates in* — See *Antikaliacs*, Section A, also *Gout*

— *Phosphatic* Acid Nitrohydrochlor Dil, and other acids Sodii Phosph. Acid in full doses, Urotropine See also *Cystitis*

— *Incontinence of* Ammon Benzoas, Acid Phosphoric, Belladonna, Buchu, Cantharis, Chloral, Creosotum, Ergota, Ferr Perchlor, Hyosciamus, Iycopodu Tinct, Potass Citras, Quinina, Sodii Benzoas, Symplicium, Iodum, Urotropine

— *Decomposing.* — See *Cystitis*

Urticaria Acid Hydrocy Dil, Acid Salicylic, Balsam Peruvianum, Ichthyol, Liquor Calcis, Potass Carb (Lotion), Sodii Salicyl, Ung Zinci, Cathartics Stomachics

Uterus, Hemorrhage of—See *Hæmorrhage*

—*Inflammation of* Acid Carbolic Glyc, Argenti Nitras, Iodoform, Iodum preps, Iodised Phenol, Zinci Sulphas

—*to contract*—See *Ecbolics*, Section A

Uvula, Inflamed Catechu Troch, Cupricum, Guaiaci Troch, Gummi Eucalyptus, Kumeri, Pyctium, Rosæ Inf Acid See also *Tonsillitis*

Vagrose Vitis Ext Ligot Liquid, Tinct Ferni Perchlor, Hamamelis

Vermifuges—Section A

Vesical Catarrh—See *Cystitis*

Vesicants—Section A

Vomiting, to allay See *Sedatives, gastric and nerve*, Section A

—*in Pregnancy* Acid Carbolic, Acid Hydrocyan Dil, Bismuth Subnit, Chloral, Chlorotone, Calcis Saccharat Liquor, Cerni Ovals, Cocaine Hydrochloridum, Chlosetum, Gentiane Inf, Ipecac, Potass Bromid, Sodii Bicarb

Warts Acid Acetic Glaciale, Acid Chromic, Acid Nitric, Argenti Nitras, Cupri Oleatis Ung, Hydrarg Nitrat Acid Liquor, Sodii Ethylatis Liquor

—*Syphilitic* Argenti Nit, Hyd Iodidi Rub Ung, Hyd Nit Acid Liquor

Wasp Sting—See *Bites and Stings of Insects*

Wax, Indurated—Glycerinum, Oleum Amygdale, Sodii Bicarb Sol

Whites—See *Leucorrhœa*

Whooping Cough Acid Acetylsalicylic, Acid Carbolic, Acid Cresylicum (initial), Acid Hydrocy Dil, Alum, Ammon Bromid, Antipyrine, Antitussin, Atropina, Bellidonna, Bromoform, Cannabis Ind, Caryoph Oleum, Chloral, Tinct Chlorofomi et Morphine Co, Conium, Eucalypti Oleum, Euphorbia Pilulif, Eugumme, Grindelia, Hydrogen Peroxid, Ipecacuanha, Lobelia, Potass Bromid, Quinina Tannas, Resorcin, Succini Lm, Trifoli Syrupus, Tussol, Zinci Sulphas

Worms, Ascarides, Tape, and Round Worms—See *Anthelmintics*, Section A

Wounds Acid Boric, Acid Carbolic, Acid Sulphuros, Acid Trichloroacetic, Aluminii Acetat Liquor, all Antiseptics (Section A), Arnica Tinctura, Tinct Benz Co, Bismuth Subnit, Collodium Flexile, Glycerinum, Iodoformum, Iodol, Sal Alenbroth

—*Poisoned* Acid Carbolic, Argenti Nitras, Hydrarg Perchlor or other antiseptic

INDEX.

The Names adopted by the British Pharmacopœia are put in Roman letters, all others whether referring to Official or Not Official Medicines, are put in Italics. The Appendix is not indexed.

As the available space for the doses in the Index is very limited, the following abbreviations are used: gr for grain, dr for drachm (either solid or fluid) or for ounce (solid or fluid), and min for minim.

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<i>Abercrombie's Cough Mixture</i>		106	<i>de Cobie bibasco</i>		465
<i>Abernethy's Pills</i>		436	<i>de Morpina</i>		782
<i>Abrastol</i>	5 to 15 gr	806	<i>de Plomb</i>		911
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<i>Cacao</i>		1203	<i>Cantharidis</i>		319
<i>Cantaridas</i>		321	<i>Cantharidina</i>		319
<i>Cotton Fylio</i>		40	<i>Ipecacuanha</i>	to to 30 min	682
<i>Fosforado</i>		884	<i>Mylabridis</i>		319
<i>de Hulecho Macho</i>		540	<i>Odonatum</i>		13
<i>de Hulecho de Bacalao</i>		790	<i>Ops</i>	8 min	841
<i>Itinaza</i>		711	<i>Sallie</i>	to to 30 min	1063
<i>Mananilla Alcanforado</i>		168	<i>Unguent</i>		1084
<i>Olivas</i>		824	<i>Acetyl methylendi diguanicol</i>		587
<i>Paranna</i>		860	<i>Acetylparamidophenol Salicylate</i>		1040
<i>Ricino</i>		1015	<i>Acetylpara amino salol</i>		1040
<i>Ruda</i>		1021	<i>Acetylphenylhydrazin</i>		5
<i>Volatil de Almonds</i>		152	<i>Acetylsalicylic Acid</i>	to to 15 gr	78
<i>Amargras</i>		398	<i>Acibar</i>		116
<i>de Canela</i>		195	<i>Barbado</i>		117
<i>Eucalypto</i>		728	<i>Sucotino</i>		118
<i>Limon</i>		765	<i>Acid values of Gums and Gum resins</i>		1313
<i>Acetanilidum</i>	1 to 3 gr	5	<i>Acid Acetique</i>		9
<i>Acetas Plumbi</i>		911	<i>crystallisable</i>		11
<i>Acetate Basique de Plomb</i>		911	<i>Azotique</i>		61
<i>Dissous</i>		919	<i>Ben oique</i>		19
<i>de Ethyle</i>		107	<i>Borgue</i>		22
<i>de Morphine</i>		782	<i>Bombyhydrigue Dissous</i>		47
<i>Neutric de Plomb</i>		911	<i>Chlorhydrigue Official</i>		50
<i>di Potassium</i>		943	<i>Chromique Crystallise</i>		37
<i>Sodium</i>		1090	<i>Citrique</i>		39
<i>Zinc</i>		1239	<i>Cyanhydrigue Dissous</i>		53

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	Dose	Page		Dose	Page
<i>Acide Gallique</i>		45	<i>Acid. F. racum</i>	2 to 5 min	41
" <i>Lactique</i>		57	" <i>Gelsemium</i>	5 to 15 gr	45
" <i>Oleique</i>		65	" "		286
" <i>Osmique</i>		67	" <i>Linum</i>		47
" <i>Oratique</i>		67	" <i>Hydrobromicum, Dilutum</i>	15 to 60 min	47
" "		68	" <i>Hydrochloricum</i>		50
" "		73	" <i>Cnidium</i>		51
" <i>Sulpureum</i>		83	" <i>Dilutum</i>	5 to 20 min	52
" <i>Sulpureum Officinal</i>		79	" <i>Hydrocyanicum Dilutum</i>	2 to 6 min	53
" <i>Tartrique</i>		89	" <i>Hydrocyanicum (Schale)</i>	1 to 5 min	55
<i>Acido Acetico</i>		9	" <i>Enthas</i>		55
" <i>Aquoso</i>		11	" <i>Hydrofluoricum</i>		55
" "		12	" <i>Dilutum</i>	5 to 20 min	56
" <i>Glacial</i>		12	" <i>Hypophosphoricum</i>	5 to 5 min	58
" <i>Hydratado</i>		10	" <i>Dilutum</i>		57
" <i>Agallio</i>		45	" <i>Iodo Salicylicum</i>		79
" <i>Arsenoso</i>		15	" <i>Kinicum</i>		394
" <i>Benzouo</i>		19	" <i>Lacticum</i>		67
" <i>Borico</i>		22	" <i>Dilutum</i>		60
" <i>Bismidrico</i>		47	" <i>Lartricum</i>		717
" <i>Camhidrico medicinal</i>		53	" <i>Myonum</i>		1079
" <i>Camhidrico</i>		53	" <i>Nitricum</i>		61
" <i>Citrico</i>		39	" <i>Cnidium</i>		62
" <i>Clorhidrico</i>		50	" <i>Dilutum</i>	5 to 20 min	64
" <i>Clorhidrico Concentrato</i>		50	" <i>Fumans</i>		62
" <i>Cromico</i>		37	" <i>Nitricum nitrosum</i>		62
" <i>Fenico</i>		27	" <i>Nitro hydrochloricum Di</i>		64
" <i>Fosforico</i>		68	" <i>lutum</i>	5 to 20 min	64
" <i>Gailico</i>		45	" <i>Nucleum</i>		809
" <i>Lactico</i>		57	" <i>Nucleum</i>		809
" <i>Lattico</i>		61	" <i>Oleum</i>		65
" <i>Nitrico</i>		61	" <i>Oleum</i>		66
" <i>Nitrico Concentrato</i>		65	" <i>Osmicum</i>		67
" <i>Oleico</i>		67	" <i>Oxalicum</i>		87
" <i>Ossalico</i>		73	" <i>a Oxynaphthoricum</i>		897
" <i>Salicico</i>		83	" <i>Paracresotinum</i>		44
" <i>Solforoso</i>		79	" <i>Paracresylicum</i>		42
" <i>Sulpureo</i>		81	" <i>Phenol para sulphonicum</i>		37
" <i>Aromatico</i>		84	" <i>Phosphoricum (conco)</i>		65
" <i>Tannico</i>		89	" <i>Dilutum</i>	5 to 20 min	70
" <i>Tartarico</i>		89	" <i>Puricum</i>		71
" <i>Tartrico</i>		89	" <i>Pyrogallum</i>		72
<i>Acidum Aceticum</i>		9	" <i>Pyrologneousum Cnidium</i>		9
" <i>Aromaticum</i>		11	" <i>Rectificatum</i>		9
" <i>Concentratum</i>	1 to 2 fl dim	11	" <i>Quillatum</i>		980
" <i>Dilutum</i>		11	" <i>Quinicum</i>		394
" <i>Glaciale</i>		11	" <i>Salicylicum</i>	5 to 20 gr	75
" <i>Acetylsalicylicum</i>		78	" <i>Sclerolum</i>		489
" <i>Agaricum</i>		112	" <i>Sphaerulicum</i>		491
" <i>Anisicum</i>		166	" <i>Stearicum</i>		461
" <i>Arsenicosum</i>		15	" <i>Sulphocarbolum</i>		37
" <i>Arseniosum</i>	1 to 15 gr	14	" <i>Sulphuricum</i>		79
" <i>Benzouicum</i>	5 to 15 gr	19	" <i>Alcoholatum</i>	5 to 20 min	82
" <i>Boricum</i>	5 to 15 gr	22	" <i>Aromaticum</i>	5 to 20 min	82
" <i>Boro-glycerinatum</i>		27	" <i>Dilutum</i>	1 to 1 fl dim	83
" <i>Cacodylicum</i>		1104	" <i>Sulphosum</i>	2 to 5 gr	84
" <i>Camphoricum</i>		313	" <i>Tannicum</i>	5 to 20 gr	89
" <i>Carbazoticum</i>		71	" <i>Tartaricum</i>		1136
" <i>Carbolicum</i>	1 to 3 gr	27	" <i>Taurocholicum</i>		11
" <i>Camphoratium</i>		36	" <i>Trihydroaceticum</i>		1229
" <i>Cnidium</i>	1 to 5 min	36	" <i>Valerianicum</i>		1230
" <i>Liquefactum</i>		33	" <i>Vanillicum</i>		417
" <i>Catharticum</i>		1076	<i>Acome</i>		91
" <i>Cetraricum</i>		358	<i>Aconiti Folia</i>		91
" <i>Chironium</i>	1 to 5 gr	394	" <i>Radix</i>		91
" <i>Chromicum</i>		37	<i>Aconitin</i>		96
" <i>Chrysophanicum</i>		39	<i>Aconitine Nitras</i>		96
" <i>Cinnamicum</i>		39	" <i>Oleatum</i>		97
" <i>Citricum</i>	5 to 20 gr	39	<i>Acondo</i>		91
" <i>Cresotinum</i>		42	<i>lequa de Calce</i>		228
" <i>Cresylicum</i>		42	" <i>Cathame</i>		909
" <i>Dibromogallium</i>		47	" <i>Distillata</i>		181
" <i>De Iodo Salicylicum</i>		79	" <i>de Arancio</i>		211
" <i>Filagicum</i>		17	" <i>Canella</i>		387
" <i>Embelicum</i>		483			
" <i>Filicicum</i>		540			

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<i>Aqua Distillata de Rose</i>		1028	<i>Aqua Femcada</i>		13
" <i>Oxygenata</i>		641	" <i>de Flores de Laranjena</i>		211
" <i>de Fiores de Alancio</i>		210	" <i>Hotela</i>		749
<i>Actææ Racemosæ Radix</i>		850	" <i>Oxygenada</i>		641
<i>Aetol</i>		187	" <i>Phenica</i>		83
<i>Adeps</i>		98	" <i>de Rabel</i>		82
" <i>Benzoinatus</i>		100	" <i>de Igele</i>		921
" <i>Induratus</i>		98	" <i>Vegeto Mineral</i>		921
" <i>lanu</i>		100	<i>Aquum</i>		1091
" <i>de Aqua</i>		102	<i>Aquum</i>		257
" <i>Hydrius</i>		102	<i>Aurogen</i>		257
" <i>de Oleo</i>		102	<i>trud</i>		277
" <i>Stallus</i>		98	<i>Asthen's Lonic Pills</i>		1006
<i>Adiposine</i>		862	<i>Atrum</i>		1
" <i>Oil</i>		860	<i>Akonit Linctus</i>		91
<i>Adiantoda</i>		102	<i>Akonit amphi</i>		640
<i>Adhesive Plaster</i>		1007	<i>Albarum</i>		177
<i>Adonidum</i>	1 to 1 gr	103	<i>Albavalde (erusa)</i>		91
<i>Adonis</i>	2 to 6 gr	103	<i>Alboferum</i>		508
<i>Adoniridera</i>		887	<i>Albumen</i>		846
<i>Adonidum</i>		1185	" <i>Ichthyolulphonas</i>		68
" <i>Chloride Solution</i>	5 to 30 min	1187	" <i>Tannus</i>		85
<i>Ailago</i>		465	<i>Alacius</i>		571
<i>Ether</i>	{ 10 to 30 min (repeated) 40 to 60 min (single) }	103	<i>Alacfor</i>		405
" <i>Aceticus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	107	" <i>Monobromada</i>		313
" <i>Alcoolizado</i>		106	<i>Alcaravea</i>		451
" <i>Bromatus</i>		109	<i>Alcarave</i>		31
" <i>Chloratus</i>		111	<i>Alcatas</i>		908
" <i>Spirituosus</i>		107	<i>Alcohol (90 p c)</i>		1146
" <i>Chloricus</i>		376	" <i>Absolutum</i>		113
" <i>cum Spiritu</i>		108	" <i>Alcanforado</i>		309
" <i>Methylatus</i>		106	" <i>Ammonia</i>		135
" <i>Nitricus Alcoholicus</i>		1140	" <i>Amlycum Testianu</i>		108
" <i>Nitrosus Spirituosus</i>		1140	" <i>Anhidro</i>		115
" <i>Ozone</i>		641	" <i>de Anus Ammoniacal</i>		166
" <i>pro Nares</i>		104	" <i>Coclearia</i>		192
" <i>Purificatus</i>		104	" <i>Cortez de Limon</i>		726
" <i>Spirituosus Camphoratus</i>		312	" <i>Dulutum</i>		1149
" <i>Sulphuricus Alcoholicus</i>		106	" <i>de Menta Piperita</i>		768
<i>Atheroleum</i>		824	" <i>Methylcum</i>	5 to 10 min	115
" <i>Citru</i>		728	" <i>de Romero</i>		1029
" <i>Laurandulu</i>		720	" <i>Table</i>		1149
" <i>Macedus</i>		799	" <i>Vinico</i>		113
" <i>Petoselem</i>		176	<i>Alcoholato de Cortez de Limon</i>		726
" <i>Rosmarini</i>		1027	" <i>Coclearia</i>		192
" <i>thorycafevum</i>	1 to 3 gr	278	" <i>Romero Compuesto</i>		109
<i>Ethyl Bromidum</i>		109	<i>Alcoholatura de Menta</i>		768
" <i>Chloridum</i>		110	<i>Alcool Absoluto</i>		115
" <i>Iodidum</i>		111	" <i>Ethylque</i>		113
<i>Ethylene Bromide</i>	1 to 2 min	109	" <i>de 95 degrees centigrade</i>		1147
" <i>Chloratum</i>		1225	<i>Alcoholatura de Anemone Pulsatilla</i>		872
<i>Ethylus Canbama</i>		109	" <i>de Citron</i>		76
<i>Ethylus Bromatum</i>		109	" <i>de Orange</i>		207
" <i>Chloratum</i>		111	<i>Alcoolatures</i>		1171
<i>Agalla de Alepo</i>		550	<i>Aldehydum Benzouum</i>		153
<i>Agallas de Levante</i>		451	<i>Alendroth Gauze</i>		626
<i>Agaricus Acid</i>		112	" <i>Wool</i>		628
" <i>Agaricus</i>	4 to 15 gr	112	<i>Aletris</i>		115
<i>Agaricus blando</i>		112	<i>Alexandrian Scana</i>		1072
" <i>Brancho</i>		112	<i>Alga Perlada</i>		559
<i>Agaricus Albus</i>	5 to 50 gr	112	<i>Alginoid Iron</i>		508
<i>Agathm</i>	5 to 10 gr	79	<i>Algoda Polvoa</i>		975
<i>Agropyrum</i>		1219	<i>Algodoero</i>		875
<i>Agua de Alcatraz</i>		909	<i>Algodon</i>		575
" <i>Alquitran</i>		909	" <i>Hydrophlo</i>		575
" <i>Brea</i>		909	<i>Alkasal</i>		129
" <i>Cal</i>		258	<i>Allyl Thiocarbamide</i>		1055
" <i>Cloro</i>		370	<i>Alnaciga</i>		780
" <i>destilada</i>		141	<i>Almendra Amarga</i>		150
" <i>de Azahar</i>		210	<i>Almendro Dulce</i>		151
" <i>de Canela</i>		397	<i>Almidon</i>		159
" <i>cortez de naranja</i>		211	<i>Almucar</i>		795
" <i>amarga</i>		211	<i>Almucle</i>		795
" <i>Alus</i>		1026	<i>Almond Pitta</i>		170
" <i>Pagedenica Negra</i>		629	" <i>Oil</i>		170
" <i>Aluja</i>		624	" <i>Sweet</i>		174
			<i>Aloe, Barbadeus</i>	2 to 5 gr	117

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	Dose	Page		Dose	Page
<i>Aloe Capensis</i>		123	<i>Ammonii Iodhylosulphonas</i>		615
" <i>Purificata</i>		119	" <i>Phosphas</i>	5 to 10 gr	147
" <i>de Socotora</i>		118	" <i>Pieras</i>	4 to 1 gr	72
" <i>Socotrina</i>	2 to 5 gr	118	" "		149
" <i>Veia</i>		117	" "		77
<i>Aloes</i>	2 to 5 gr	116	" <i>Valerianas</i>		149
" <i>de Barbades</i>		117	<i>Ammonio Chloride of Mercury</i>		671
<i>Alomum</i>	1 to 2 gr	120	" <i>Fine Alum</i>		534
<i>Alpha Egon</i>	5 to 10 gr	847	" <i>Mercuric Chloride</i>		625
" <i>Naphthol</i>		804	<i>Ammonio</i>		132
<i>Alphol</i>		807	" "		121
<i>Alquistan</i>		908	" "		144
<i>Alsol</i>		125	" <i>Hydricum Solutum</i>		132
<i>Alstoma</i>		124	" <i>Jodatum</i>		146
<i>Altea</i>		124	" <i>Sulphate</i>	5 to 15 gr	149
<i>Althæa Radix</i>		124	" <i>Sulphate of Potash</i>		7
<i>Alum Rose Gargle</i>		127	" <i>Sulphate of Soda</i>		637
<i>Alumen</i>	5 to 10 gr	125	" <i>Valerianate</i>	1 to 3 gr	149
" <i>Extinctum</i>		127	" <i>Valerianatum Solutum</i>		149
" <i>Ustum</i>		127	" "		136
<i>Alumini Acetas Solutio</i>		127	" "		127
" <i>Aceto Tartar</i>		128	<i>Ammoniumcarbonat</i>		139
" <i>et Ammonii Sulphatas</i>		129	<i>Ammoniumchlorid</i>		144
" " <i>Sulphas</i>		125	<i>Ammonol</i>	5 to 10 gr	5
" <i>Borotartar</i>		129	" <i>Bromide</i>		5
" <i>Borotannas</i>		129	" <i>Lithate</i>		5
" <i>Caseinas</i>	5 gr	129	" <i>Sulphate</i>		5
" <i>Chloridum</i>	5 to 8 gr	128	<i>Ammonio</i>		122
" <i>Hydricum</i>		128	<i>Amoras</i>		778
" <i>Naphthol Sulphonas</i>	4 to 8 gr	128	<i>Amygdala Amara</i>		150
" <i>Nitras</i>		128	" <i>Dulcis</i>		154
" <i>Oleas</i>		128	<i>Amalgam (Mercurial)</i>	1 to 4 fl drim	140
" <i>et Potassii Sulphatas</i>		120	<i>Amertatin</i>		969
" <i>Sulphas</i>		120	<i>Amigdophenina</i>	8 to 15 gr	875
" <i>Sulphas</i>		129	<i>Amyl Nitris</i>	2 to 5 min (in dilution)	155
" <i>Sulphas</i>		128	" <i>Tertius</i>	5 drops	155
<i>Alumol</i>		128	" <i>Sulphas</i>		535
" "		125	" <i>Valerianas</i>	2 to 3 min	138
" "		914	<i>Amylene Chloral</i>		366
<i>Alupin</i>	1 to 1 gr	417	" <i>Hydrate</i>	50 to 60 min	138
<i>Amandes Amarae</i>		150	<i>Amplenium Hydricum</i>		135
" <i>Doses</i>		154	<i>Amylic Alcohol Tincture</i>		135
<i>Anapola</i>		1017	<i>Amyloform</i>		146
<i>Ambar</i>		1172	<i>Amyloform</i>		146
" <i>Anardillo</i>		1172	<i>Amylum</i>		159
<i>Amber</i>		1172	" <i>Jodatum</i>		161
" <i>Oil of</i>		1172	<i>Anaesthesia</i>	5 to 10 gr	23
<i>Amezas Passadas</i>		971	" "		23
<i>Amendos Amargas</i>		150	" "		375
" <i>Doses</i>		154	<i>Anerthyl</i>		111
<i>Amerikanische Kreuzdominde</i>		337	<i>Analgia</i>	7 to 15 gr	101
<i>Amidon de Ble</i>		159	<i>Anarcotina</i>		844
<i>Anime Acid</i>		44	<i>Andee's Lotion</i>		1010
<i>Aninaform</i>		515	<i>Andioaphus</i>		481
<i>Ammonia Liquida</i>		132	<i>Ammonia Camphor</i>		972
<i>Ammonia</i>		132	<i>Aneth</i>		162
<i>Ammoniacum</i>	5 to 15 gr	129	<i>Anethi fructus</i>		162
<i>Ammoniakum</i>		129	<i>Anethol</i>		166
<i>Ammoniaque Official</i>		132	<i>Aneto</i>		162
" <i>Dilute</i>		133	<i>Angostura Verduleria</i>		470
<i>Ammoniacal Tincture</i>		135	<i>Angostura Arunde</i>		489
" <i>Muciv</i>		631	<i>Angusture Franc</i>		489
" <i>Tincture of Ergot</i>		488	<i>Angusturin</i>		469
" " <i>Guaiacum</i>		588	<i>Anhydride Arsenic</i>		11
" " <i>Opium</i>		841	<i>Anhydride Gluco Chloral</i>		368
" " <i>Quinine</i>		1005	<i>Anice</i>		163
" " <i>Vile lau</i>		1228	<i>Anidride Arsenosa</i>		14
<i>Ammonii Benzoas</i>	5 to 15 gr	136	" <i>Cromua</i>		37
" <i>Boarboas</i>		144	<i>Aniline</i>		163
" <i>Boras</i>		137	<i>Animal Charcoal</i>		326
" <i>Bromidum</i>	5 to 30 gr	137	<i>Amas Commu</i>		165
" " <i>Pfervescens</i>		139	" <i>Fert</i>		163
" <i>Carbonas</i>	3 to 10 gr	139	<i>Aniseed Cordial</i>		166
" <i>Chloridum</i>	5 to 20 gr	144	<i>Anisi Fructus</i>		163
" <i>Embelas</i>	3 to 6 gr	483	<i>Anise Acid</i>		166
" <i>Fluoridum</i>		56	<i>Anissamine</i>		163
" <i>Iodidum</i>	2 to 5 gr	146	<i>Anodyne Collodion</i>		977

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ARE

	Dose	P.L.		Dose	P.L.
<i>Anodyne Collod</i>		977	<i>Aponomphum Hydric</i>	{ 1 to 1 fl oz (hypodermic) 1 to 1 fl oz (oral)	178
" <i>Spirit Hoffmanni</i>		106	<i>chloridum</i>		
<i>Anthemidis Flores</i>		167	<i>Amorphine, Dosis of</i>		180
<i>Anthrapurpurin Diacetate</i>		1017	<i>Apozem</i>		859
<i>Anthracobin</i>		180	" <i>de Cousin</i>		471
<i>Anti Anthrax Serum</i>	1 to 2	16	" <i>Grenadier</i>		675
<i>Antibubonic Pills, Little</i>		12	" <i>Purgatif</i>		1075
<i>Anti catarrhal Salts</i>		31	<i>Applatio Menthol</i>		771
<i>Antidotes</i> See under respect			" <i>Sinapis</i>		1044
live herding			<i>Aqua Ammonia</i>		183
<i>Antidatum Ischnus</i>		15	" <i>Portion</i>		182
<i>Antijebria</i>		5	" <i>Amigdala Amara</i>		182
<i>Antikamnia</i>		8	" <i>Anethi</i>	1 to 1 fl oz	162
<i>Antimonial Powder</i>		170	<i>Anisi</i>	1 to 1 fl oz	164
" <i>Wine</i>		173	" <i>Anthemidis</i>		168
<i>Antimonii Oxidum</i>	1 to 2	16	" <i>Amantii Floris</i>		210
" <i>et Potassii Tartras</i>		171	" <i>Calcanis</i>		288
<i>Antimonotartrate Acidi de</i>			" <i>Calis</i>		288
<i>Potassium</i>		173	" <i>Saccharata</i>		288
<i>Antimonious Oxide</i>		169	" <i>Camphor</i>	1 to 2 oz	307
" <i>Sulphidi</i>		171	" <i>Conc</i>		311
<i>Antimonium Nigrum Purificatum</i>		171	" <i>Citobolus</i>		33
" <i>Crude</i>		171	" <i>Carbolisata</i>		35
" <i>Sulphuratum</i>	1 to 2 gr	171	" <i>Cinn</i>	1 to 2 fl oz	331
" <i>Tartratum</i>	1 to 2 gr	173	" <i>Chlorata</i>		370
<i>Antinosm</i>		882	" <i>Chlori</i>		369
<i>Antipyrina</i>		477	" <i>Chloroformi</i>	1 to 1 fl oz	375
<i>Anti Plague Serum</i>		1263	" <i>Cinnamomi</i>	1 to 2 fl oz	396
<i>Anti Pneumococci Serum</i>		1263	" <i>Spirituosa</i>		397
<i>Antipyronine</i>		26	" <i>Cicosta</i>		454
<i>Antipyrium</i>		877	" <i>Cicostia</i>		43
<i>Antipyrima ferussens</i>		479	" <i>Destillata</i>		181
<i>Antipyrine</i>		877	" <i>Emuncti</i>	1 to 2 fl oz	541
" <i>Aceto Salicylat</i>		879	" <i>Goulardi</i>		921
" <i>Amigdalo</i>		881	" <i>Hydrogenii Dioxide</i>		641
" <i>Salicylate</i>		880	" <i>Acresolita</i>		42
<i>Antipyrinum Caffeina utrum</i>		880	" <i>Immutata</i>	1 to 2 fl dim	718
<i>Anti Rabii Inoculation</i>		1268	" <i>Lithia (Lithrocarbonis) See</i>		
<i>Antirheumatis</i>	1 to 2 gr	774	" <i>Iquor Lithia Carbonatis</i>		735
<i>Antirheumatis</i>		774	" <i>Mellis</i>		764
<i>Antiscorbuto Syrup</i>		192	" <i>Meuth Piparito</i>	1 to 2 fl oz	767
<i>Antiseptum</i>		8	" <i>Vindis</i>	1 to 2 fl oz	769
<i>Antiseptic Dressing, Lora</i>		26	" <i>Menthol</i>		771
" <i>Carbolic</i>		5	" <i>Naphae</i>		210
" <i>Bucaliphys</i>		405	" <i>Opi</i>		841
" <i>Iodoform</i>		664	" <i>Petroclini</i>		176
" <i>Salicylic</i>		77	" <i>Fenicata</i>		33
" <i>Thymol</i>		1210	" <i>Phenolata</i>		33
" <i>Mouth Wash</i>		35	" <i>Puis</i>	1 to 2 pints	909
" <i>Solution</i>		1154	" <i>Piment</i>	1 to 2 fl oz	899
<i>Antiseptin</i>		1251	" <i>Plumbi</i>		921
<i>Antiseptol</i>		394	" <i>Goulardi</i>		921
<i>Anti Sera</i>		125	" <i>Spirituosa</i>		921
<i>Antispasmin</i>	1 to 2 gr	844	" <i>Plumbica</i>		921
<i>Anti Streptococci Serum</i>		1202	" <i>Picroli Pini</i>		909
<i>Antitoxin</i>		8	" <i>Stabelli</i>		82
<i>Antitoxins</i>		1260	" <i>Iegua</i>		65
<i>Anti Tubercle Serum</i>		1263	" <i>Rose</i>		1026
<i>Antivirusum</i>		56	" <i>Portion</i>		1026
<i>Anti Typhoid Extract of Jc</i>		1263	" <i>Sinibuci</i>		1041
" <i>Serum</i>		1263	" <i>Sativa</i>		921
<i>Antivenene</i>		1262	" <i>Alcoolisada</i>		921
<i>Antivenomous Sera</i>		1262	" <i>de Vegeto</i>		920
<i>Antrophores</i>		1202	" <i>Vegeto mineralis Goulardi</i>		920
<i>Anusol</i>	2 to 5 min	1010	<i>Aqua (group)</i>		181
<i>Anytin</i>		658	<i>Arabisches Gummi</i>		1
<i>Anytol</i>		658	<i>Arachis Oil</i>		827
<i>A Oxyanaphthoic Acid</i>		807	<i>Arancio Amaro</i>		206
<i>Aphthura</i>		587	<i>Asatoba</i>		183
<i>Apol</i>		176	" <i>Depurata</i>		379
" <i>Cryst</i>		177	<i>Araruta</i>		760
<i>Apocodena</i>		426	<i>Arbutin</i>	1 to 15 gr	1226
<i>Apocodena Hydrochloridum</i>		426	<i>Arena</i>		183
<i>Apocynin</i>		177	" <i>Nut Charcoal</i>		184
<i>Apocynum</i>		177	<i>Arecolinae Hydrobromidum</i>	1 to 1 gr	184
<i>Apolyvin</i>		876			

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	Dose	Page	
		187	<i>Aspidosamine</i>
		188	" <i>Hydrochloride</i>
" <i>Curas</i>		188	" <i>Hydrochloride</i>
" <i>Fluoridum</i>		189	" <i>Hydrochloride</i>
" <i>Ichthyolas</i>		188	<i>Aspirin</i>
" <i>Iodidum</i>		187	<i>Assenzio</i>
" " <i>Nascens</i>		187	<i>Asucar</i>
" <i>Lactas</i>		187	" <i>de Leite</i>
" <i>Nitras</i>	$\frac{1}{2}$ to $\frac{1}{2}$ gr	184	<i>Asterol</i>
" " <i>Fusus</i>		186	Atomic Weights and Symbols
" " <i>Induratus</i>		186	<i>Atozol</i>
" " <i>Mitigatus</i>		186	<i>Atropina</i>
" <i>Nitric Styl</i>		186	" " "
" <i>Nucleinas</i>		810	" " "
" <i>Ocidum</i>	$\frac{1}{2}$ to 2 gr	190	" <i>Salicylas</i>
<i>Argentol</i>		187	" <i>Sulphus</i>
<i>Argentum Foliatum</i>		187	" " "
" <i>Nitricum c. Kalio Nitrico</i>		186	" " "
<i>Argonin</i>		187	<i>Atropinsulfat</i>
<i>Argonin L</i>		187	<i>Atroscine</i>
<i>Argyrol</i>		187	" <i>Hydrobromide</i>
<i>Arhovin</i>	1211		" " "
<i>Aristochin</i>	983		" " "
<i>Aristol</i>	1211		" " "
<i>Aristolochia</i>	1077		" " "
<i>Almoracin Radix</i>	191		" " "
<i>Alnica Opodeldoc</i>	193		" " "
<i>Alnica Flores</i>	193		" <i>Niles Cortex</i>
" <i>Rhizoma</i>	192		" <i>Bromonium</i>
<i>Alnikavurzel</i>	192		" <i>Chloridum</i>
<i>Aromatic Acetic Acid</i>	13		" <i>et Potassium Bromidum</i>
" <i>Cascara</i>	340		" <i>Sodium Chloridum</i>
" <i>Infusum of Cassia</i>	563		" " "
" <i>Infusum of Cassia</i>	457		" " "
" <i>Infusum of Cassia</i>	141		<i>Azonges</i>
" <i>Syrup</i>	81		<i>Azungra</i>
" <i>Vinegar</i>	206		" <i>Porca</i>
<i>Arrhenal</i>	13		<i>Azadiraachta Indua</i>
<i>Arrova</i>	1105		<i>Azahan</i>
<i>Arrow Root</i>	760		<i>Azeite</i>
<i>Arseni Tricloridum</i>	760		<i>Azotate d'Argent</i>
<i>Arsenate de Fer</i>	15		" <i>de Broyde de Mercure</i>
" <i>Sodium Officinale</i>	508		" <i>Dissous</i>
<i>Arsenate de Fierro</i>	1092		" <i>Neutre de Bismuth</i>
" <i>Hierro</i>	509		" <i>de Potassium</i>
" <i>Bisodico</i>	508		" <i>Potassium</i>
" <i>Sodico</i>	1092		<i>Azotate d'Amyle</i>
<i>Arsenic</i>	1002		<i>Azucar</i>
<i>Arsenical Caustic Powders</i>	14		" <i>de Cana</i>
" <i>Paste for Dentists</i>	19		" <i>Leche</i>
" <i>Paste for cancer (Fries)</i>	19		<i>Azufre</i>
" <i>Cones</i>	19		" <i>Dorado de Antimonio</i>
" <i>Paste for Dentists</i>	19		" <i>Precipitado</i>
" <i>Solution</i>	17		" <i>Sublimado y Lavado</i>
<i>Arsenic Antidotum</i>	15		<i>Azul de Metilena</i>
<i>Arsenige Saure</i>	14		
<i>Arsenii Bromidi Liquor</i>	194		<i>Bael Fruit</i>
" <i>Iodidum</i>	$\frac{1}{2}$ to $\frac{1}{2}$ gr	194	<i>Baile's Pill</i>
<i>Arsenious Acid</i>	14		<i>Bain Alcalin</i>
" <i>Anhydride</i>	14		" <i>Sulfure</i>
<i>Arsenicaures Eisen oxydul</i>	508		" <i>Liquide</i>
<i>Arnica</i>	1105		<i>Baird's Pill</i>
<i>Artemisan</i>	1046		<i>Baldrian</i>
<i>Artemann's Creolin</i>	43		<i>Balneum Alkalinum</i>
<i>Asafetida</i>	196		" <i>Sulphuretum</i>
<i>Asant</i>	196		<i>Balsamo de Copayre</i>
<i>Asaprol</i>	$\frac{1}{2}$ to $\frac{1}{2}$ gr	806	" <i>Opodeldoch Liquido</i>
<i>Asepsin</i>	8		" <i>Solido</i>
<i>Aseptic Shaving Cream</i>	863		" <i>Peruvianum</i>
" <i>Wax</i>	352		" <i>Tolutano</i>
<i>Aseptin</i>	26		" " "
<i>Aseptol</i>	87		" " "
<i>Aseptoline</i>	896		" " "
<i>Asiatic Pills</i>	18		" <i>Peruvianum</i>
<i>Asparagin</i>	125		" <i>Tolutanum</i>
<i>Aspidium</i>	539		" <i>Tranquilli</i>

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Banika		98	" " hydroxy Naphthalene		804
Baptisin	1 to 5 gr	221	Beta naphthol		804
Bathados Aloes		117	" camphathyl orthoxy meta tokylic Acid		807
Barbalum		120	Betel	2 to 8 gr	242
Barentraubenblätter	1 to 2 gr	1226	Betol		806
Bara Chloridum		221	Bettendorf's Test		1816
" Sulphurum		221	Betulae Albe Oleum		242
Barley, Pearl		601	Beurre de Cacao		120
Basilicon Ointment		1005	" Muscade		800
Basische Natriumthymatrat		250	Bibergel		346
Bassorin		1218	Pilorate of Sodium		259
Battley's Liquor Opi Sedativus		842	Lican bonas Kalcus		936
Bauine Opodiloch		1053	" Potassu		936
" du Perou		214	Bicar bonate di Potassio		935
" de Poli		217	Lican bonato de Potassa		936
" Tranquille		648	" di Sodio		1096
" Yie (Decoct Aloes Co)		121	Bichloride of Mercury		621
Bay Khum		900	Bichloridato de Chinnu		994
Bazin's Ointment		609	Bichlorure de Mercur		621
Beberberia Leaves	1 to 5 gr	1226	Bichloruro di Mercurio		621
Beberina Sulphas		222	Bromato di Ictassio		937
Beberine	1 to 2 gr	222	Ligan ader		206
" Hydrochloride	1 to 2 gr	222	Biodure de Mercur		610
" Sulphate		222	Bismkrauththalen		644
Beberu Bark		222	Bismode of Mercury		610
Beer Yeast		1236	Bogen		845
Beeswax, White		352	Brydure di Mercurio		610
" Yellow		849	Bryde de Manganuse		757
Bela Fructus		222	Buch Tar Oil		242
Belene		644	Buch, Oil of Sweet		553
" Negro		644	Budan	1 to 4 gr	795
Belladonnablätter		224	Bismal		257
Belladonna Folii		224	Bismone		254
" Radix		228	Bismuth Cyan		240
Belladonnaum-el		228	Bismuth Phenol	5 to 20 gr	256
Benecina		241	" Purip		243
Benjoin		237	" Tetra viodophenolphthalen		832
Benzaldehydum		152	" Tridromophenol		258
Benzalgen		101	Bismuthi et Ammonii Citras	2 to 5 gr	250
Benzin		237	" " " Effer vesens		254
Benzina		241	" Benzoes	5 to 20 gr	258
Benzinum Purumatum		237	" Beta Naphtholas	5 to 20 gr	243
Benzosadricus cum Coffeino		270	" Carbas	5 to 20 gr	249
" Natrio cofficus		276	" Ceru Salicylas	5 gr	258
Benzonate d Ammonium		130	" Cinchonidine Iodidum		1111
" de Sodium		1094	" Cinchamas	2 to 5 gr	255
Benzonitrid Irid		100	" Citras		258
Benzonate di Ammonio		130	" Di tho Salicylas		258
" Lituo		757	" Iodo Kesorin Sulphonas		257
" " effervescenti		737	" Methylenedigallas		261
" de Lituo		737	" Nitras		256
" " Sodio		1094	" Oleas	5 to 20 gr	247
" " e Caffeeia		276	" Oxidum		247
Ben oharz		19	" " (Colloidum)		247
benzoazure		504	" " Hydritum		247
benzoazureumidum		19	" Oxycar bonas		255
Benzoic Acid		22	" Oxychloridum		257
" " Torenge		22	" Oxynodogallas		250
Benzoium		237	" Oxynitias		246
Benzoil	3 to 5 min	241	" Phenias		258
Benzonaphthol	5 to 15 gr	508	" Phosphas		258
Benzosol		585	" Quinolun		258
Benzosulphuridum		504	" " " Sulphocyanum		258
Benzoyl Morphine Hydrochloride	1 to 4 gr	750	" Salicylas	5 to 20 gr	247
" Peroxide		845	" Subgallas		250
" Pseudoto opavine		416	" Subiodidum	5 to 10 gr	258
" Hydrochloride		416	" Subnitras	5 to 20 gr	250
" Sulphonimide		564	" Sulphas		255
Berberina Phosphas	1 to 5 gr	242	" Tinnas		257
Berberine		241	" Tridromophenolas		255
Berberis		105	Bismuthum		243
Bertonis Ethel		105	Bismuto		243
Betula		105	" Carbonato		243

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<i>Bitartras Kalicus</i>		966	" <i>de Potassio</i>		939
" <i>Potassia depuratus</i>		966	<i>Brometum Ethylicum</i>		109
Bitartrate of Potassium		966	" <i>Kalicum</i>		939
<i>Bitartrato de Potassa</i>		966	<i>Bromhidrato di Scopolamina</i>		649
" <i>Potassio</i>		966	<i>Bromhidrato d Homatropina</i>		598
<i>Bittere Mandeln</i>		150	" <i>d Hyoscyne</i>		649
Bittersweet		480	" <i>de Quinine Nucle</i>		966
<i>Black Cat's Claw</i>		345	<i>Bromidia</i>		366
" <i>Black</i>		1074	<i>Bromidato di Omatropina</i>		598
" <i>Black</i>		1233	<i>Bromine</i>		262
" <i>Mercurial Lotion</i>		629	<i>Bromopin</i>		261
" <i>Oxide of Manganese</i>		756	<i>Bromo</i>		262
" <i>Pepper</i>		902	"	3 to 15 gr	8
" <i>Pitch</i>		910	"	" to 3 min	8
" <i>Wash</i>		639	<i>Bromoforn</i>		26
" <i>Wattle</i>		4	<i>Bromohermol</i>		593
<i>Bladder Wrack</i>		515	<i>Bromol</i>		36
<i>Blanc de Baleine</i>		155	"		850
<i>Blueholz</i>		591	" <i>de Potassio</i>		675
<i>Blaud's Pills</i>		513	" <i>de Potassio</i>		55
<i>Bleuetat</i>		910	<i>Bromum</i>		261
<i>Bleuesig</i>		919	<i>Bromural</i>	5 to 10 gr	1225
<i>Bleiglatte</i>		916	<i>Bromine d Ammonium</i>		117
<i>Bleijodid</i>		915	" <i>d Ethyle</i>		109
<i>Blennostasine</i>		993	" <i>de Potassium</i>		939
<i>Bleu de Méthylene officinali</i>		774	" <i>Sodium</i>		1100
Blistering Collodion		320	" <i>Silicium</i>		1157
" <i>Liquid</i>		320	<i>Bromietum Kalici</i>		939
<i>Blue Cohosh</i>		349	" <i>Potassii</i>		939
" <i>Pill</i>		604	<i>Bromino di Ammonio</i>		117
<i>Blutegel</i>		597	" <i>Etile</i>		100
<i>Boa d of Health Cholera Mixture</i>		457	" <i>Estomaco</i>		1157
<i>Bodelha</i>		548	" <i>Ferrug</i>		510
" <i>Boa d of Health</i>		591	" <i>de Luto</i>		768
<i>Boldo</i>		258	" <i>Oro</i>		212
<i>Boldo</i>		258	" <i>Potassio</i>		939
<i>Bolus Alba</i>		709	" <i>de Potassio</i>		939
<i>Bometero</i>		499	" <i>di Sodio</i>		1100
<i>Bone Black</i>		526	" <i>Silicio</i>		1157
" <i>Marrow, Red</i>		761	"		47
<i>Boneset</i>		501	<i>Brown Squard's Oculitis Fluid</i>		1066
<i>Bons Blaster</i>		322	<i>Bucine</i>	1 to 1 gr	1139
<i>Boracic Acid</i>		2	" <i>Nitrate</i>		823
<i>Boral</i>		129	" <i>Sulphate</i>		821
<i>Borate de Soude</i>		259	<i>Bryonia</i>		265
<i>Borato Sodico</i>		259	<i>Buco</i>		265
<i>Borax</i>	5 to 20 gr	259	<i>Bucoblatter</i>		265
<i>Boile Acid</i>		22	<i>Buchi Bolia</i>		265
" <i>Ganze</i>		26	<i>Buckbean</i>		773
" <i>Lent</i>		26	<i>Bugnarum Zinc Sulphatis</i>		1251
" <i>Ointment</i>		25	<i>Bulbo de Colchico</i>		426
" <i>Wool</i>		26	" <i>Colchico</i>		426
<i>Borneol</i>		305	<i>Bursandy Pitch</i>		906
" <i>Sakaylate</i>		305	<i>Burrow's Solution</i>		125
" <i>Valerianate</i>		305	<i>Buro di Cacao</i>		1203
"		305	<i>Busselole</i>		1226
" <i>Potassium</i>		968	<i>Butea</i>		711
<i>Borsaine</i>		22	<i>Butea Gummi</i>		711
<i>Bouges, Iodoform and Eucal</i>		603	" <i>Semina</i>		711
<i>typtus</i>		603	<i>Butternut</i>		706
<i>Boulton's Solution</i>		671	<i>Butea</i>		867
<i>Bourbous Water</i>		19	<i>Butyl Chloral Hydras</i>	5 to 20 gr	267
"		26	<i>Byne</i>		268
" <i>Solution</i>	1 to 2 fl oz	931	<i>Lynan</i>		270
<i>Brandy</i>		1150	<i>Cabalonga</i>		658
<i>Brazil Wood Solution</i>		109	<i>Cacao Butter</i>		1203
<i>Brea</i>		908	<i>Cachou</i>		346
" <i>de Orvedio</i>		271	" <i>de Pingu</i>		346
<i>Brechnuss</i>		510	<i>Cacodylic Acid</i>		1104
<i>Brechweinstein</i>		173	<i>Cade Bath Fluid</i>		271
<i>Brechwurzel</i>		674	<i>Calamin Oleni</i>		271
<i>Brodie's Gout Pill</i>		422	<i>Calamin Sulphur</i>		1251
<i>Bromaline</i>		264	<i>Calamin Iodide</i>		919
<i>Brome</i>		262			
<i>Bromethylform</i>	5 to 30 gr	764			

CAR	Official Names in Roman, all others in Italics				
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Carbo Animalis		326	Cassa Bark		492
" Ligni	60 to 120 gr	326	Cassena Saggiada		337
Carbol. Lysoformi		547	" " " "		339
Carbolic Acid		27	" " " "		340
" " Crudi		36	Cascargilia		341
" " Dissangi		35	Cassanilla		341
" " Gauze		35	Cassanilla		341
" " Ligatures		35	Cassanilla		341
" " Lint		35	Cassia Beatedna		344
" " Liquefied		33	Cassia Oleum	1 to 5 min	343
" " Mouth Wash		35	" Pulpa	60 to 120 gr	345
" " Protective		35	Castile Soap		1043
" " Sutures		35	Castor Oil		1015
" " Tow		35	Castoreo		346
" " Wool		35	Castoreum	5 to 10 gr	346
Carbolic Soaps		35	Cataplasma Kaulum		709
" Oil		28	" Lani		732
" Solution		26	" Sinaps		1081
Cubon Disulphide		327	Catichum	5 to 15 gr	346
" Tetrachloride		377	" Nigron	5 to 15 gr	345
" Vegetal		327	" Pallidum		346
" " Medusinal		326	Catern		346
Carbonas Ferri Saccharatus		511	Cathartic Acid		1076
" Kalous		943	Cathartic, Oil for		35
" Plumbeus		914	Cato		347
Carbonate de Lithine		732	Catrame Vegetale		908
" " Magnesium		750	Cauceru		322
" " Plomb		913	Caucho		322
" " Zinc		1241	Caulo " "		349
" Acide de Sodium		1006	" " " " Hydrochloride		349
" Neutre " " Crystalliz			" " " " Mild	1 to 4 gr	349
" " " " Official		1107	" " " " Potash		187
" (resqin) d'Ammoniaque		139	" " Soda		1086
" Acide de Potassium		935	Caustico di Vienna		931
" Neutre " " "		942	" " " " "		671
Carbonato di Ammonio		139	" " " " "		1244
" de Bismuto		243	Cebaea		601
" " Cal		280	Cedarine		546
" de Calcio		280	Cedro		722
" de Chaux Precipiti		280	Celandine		360
" " Pierro		511	Celloidin		977
" di Litina effervescente		733	Cephaeline		692
" de Litina o Lituo		732	" Hydrochloride		602
" " Litio		732	Cera Alba		353
" " Manganeso		757	" Amasella		350
" di Magnesio		748	" Amarilla		349
" de Plomo		914	" Blanca		352
" Potasio		942	" Blanca		353
" de Potasio acido		936	" Blanca		349
" " Neutro		943	" Flava		349
" di Potasio		942	" Gialla		1026
" Sodico Crystallizado		1107	Cerat à la Rose		1026
" de Sodio		1107	" de Galen		358
" Zincuo		1241	Cerato de Galeno		353
Carbonis Bisulphidum		327	" Simple		353
Carbonylamide		1238	Ceratum		353
Cardamomes		328	" Calammar		280
Cardamomi Semina		328	" Camphore		311
Cardamomo		328	" Cantharidis		321
" minor		328	" Cetacei		358
Cardenillo		466	" "		363
Carlsbad Salt, Effervescent			" "		921
" Powder of		1132	" Resine		921
" Armine		421	" " Compositum		1008
Carinferrin		508	Cereoli		1008
Caripaine		855	Ceru Ovalas	2 to 10 gr	1202
" Hydrochloride		855	Cerum Oxalicum		353
Cariragen		359	Ceroto Simple		353
Carion Oil		289	Cerussa		913
Carta Senapata		1052	Cetaceum		355
Carui Fructus		331	Cetina		355
Carvacrol Iodide		1211	Cetraria		355
Carui		331	Cetraria Acid		358
Carvone		333	Cetrarin		358
Carvonin		333	Cevada Santa		358
Caryophyllum	5 to 10 gr	333			601

Official Names in Roman, all others in Italics			CHO		
	Dose	Page		Dose	Page
<i>Cevadilene</i>		1232	<i>Chloralum Hydratum</i>		362
<i>Cevadine</i>		1231	<i>Chloras Kalcus</i>		945
<i>Cevadurum</i>		1232	<i>Chlorate de Potassium</i>		944
<i>Characilla</i>		341	<i>Chlor butyl Alcohol</i>		369
<i>Chalk French</i>		1192	<i>Chloreto de Ammonio</i>		145
" Mixture		456	" " <i>Calco</i>		283
" Precipitated		280	" <i>Peruo Anhydro</i>		521
" Prepared		456	" <i>Crystallizado</i>		521
<i>Chamomile Flowers</i>		167	" <i>Mercurio</i>		623
<i>Chanori, Indian</i>		314	" <i>Mercurio</i>		627
<i>Charcoal Vegetal Official</i>		326	" <i>de Olio</i>		212
<i>Charcoal Animal</i>		326	" <i>de Sodio</i>		212
" <i>Isocids</i>		27	<i>Chlorotoni</i>	5 to 20 gr	369
" <i>Caprules</i>		327	" <i>Phru</i>		369
Wood		326	<i>Chlorotum Ethylicum</i>		111
<i>Cratichnepestus</i>		322	" <i>Amido hydrargyricum</i>		631
" <i>Nitrate</i>		959	" <i>Ammonium</i>		144
" <i>et Chlorate</i>		959	" <i>Chimicum</i>		982
" <i>Sinapius</i>		1082	" <i>Pericum</i>		521
" <i>Sinapis</i>		1082	" <i>Hydrargyricum Ammonium</i>		631
" <i>Sinapicata</i>		1082	" <i>Hydrargyricum</i>		621
<i>Chaulmoogra Oil</i>	5 to 10 min	590	" <i>Carosivum</i>		623
<i>Chaulmoogra Acid</i>		590	" <i>Hydrargyricum</i>		627
<i>Chelidonium</i>	1 to 5 gr	360	" " <i>Precipitatum</i>		627
" <i>Hydrochloride</i>		360	" " <i>ope Vaporis Aqua</i>		627
" <i>Sulphate</i>		360	" <i>Pilocarpicum</i>		896
" <i>Tannate</i>		360	<i>Chlorhydras Quininae</i>		992
<i>Chelidonium</i>		360	<i>Chlorhydrate d Apomorphine</i>		178
<i>Chelsen Pensoni</i>	583	1179	" <i>de Cocaine</i>		405
<i>Chemical Food (Squills)</i>		530	" <i>Morphine</i>		784
<i>Chene</i>		979	" <i>Lavique de Quinine</i>		991
<i>Cherry Laurel Leaves</i>		717	" <i>neutrie de Quinine</i>		994
" " <i>Witch</i>		718	<i>Chloro Liquor</i>	10 to 20 min	369
<i>Chenopodium</i>		663	<i>Chloro Ether</i>		376
<i>Chenopodium</i>		1198	<i>Chloride de Apomorphine</i>		175
<i>Chenopodium</i>		1220	" " <i>Chimica</i>		991
<i>Chile</i>		521	" " <i>Morphine</i>		784
<i>China Calisaya</i>		582	<i>Chlorinated Lime</i>		298
" <i>Camphor Oil</i>		312	<i>Chlorine Solution of</i>		369
" <i>Clay</i>		709	<i>Chloralk</i>		298
" <i>Rosa</i>		382	<i>Chlorobrom</i>	½ to 1 fl oz	368
" <i>Succubra</i>		382	" <i>Chlorodyne</i>		376
<i>China Fluide Extractum cum</i>		388	" <i>Loenges</i>		377
" <i>Kalio Iodati</i>		392	<i>Chlorodynum</i>		376
" <i>Vinum</i>		392	<i>Chloroform pro narcosis</i>		373
<i>Chinaphenun</i>	5 to 15 gr	576	<i>Chloroforme Anæsthesique</i>		373
<i>Chinarinde</i>		382	" <i>rectifié du commerce</i>		373
<i>Chinesischer Summt (Cassia)</i>		395	<i>Chloroformum</i>	1 to 5 min	371
<i>China Acid</i>		394	" <i>Aconiti</i>		95
<i>Chinina</i>		981	" <i>Belladonna</i>		235
<i>Chininum bisulphas</i>		988	" <i>Camphoratum</i>		377
" <i>bisulphuricum</i>		988	" <i>Hyoscyama</i>		647
" <i>Ferro Citricum</i>		516	" <i>Iodi</i>		671
" <i>Hydrochloricum</i>		992	<i>Chlorum Solutum</i>		370
" <i>Sulphuricum</i>		999	<i>Chlorure d Ammonium</i>		144
<i>Chinoidin</i>		990	" <i>de Chaux</i>		298
<i>Chinolin</i>		990	" " <i>Calco</i>		282
" <i>Periodide</i>		991	" " <i>Crystallus</i>		283
<i>Chinosol</i>		547	" " <i>Fondu</i>		283
<i>Chinotropane</i>	10 to 15 gr	360	" <i>d Ethyle</i>		110
<i>Chirata</i>		360	" <i>de Methyle</i>		773
<i>Churette</i>		375	" <i>d Or et de Sodium</i>		212
<i>Chloroethoform</i>		110	" <i>de Sodium Official</i>		1109
<i>Chloroethyl</i>		368	" <i>de Soude Dissous</i>		301
<i>Chloral Camphoratam</i>		368	" <i>Lime</i>		1242
" <i>cum Cocaine</i>		368	<i>Chloruro Ferrico</i>		521
" <i>Carbolatum</i>		367	" <i>Mercurio</i>		626
" <i>Formamide</i>		367	" <i>Sodico</i>		1109
" <i>Hydras</i>	5 to 20 gr	362	<i>Chlor Zinc Iodine (Schulze's)</i>		1244
" <i>et Phenol</i>		368	" <i>Solutions</i>		1244
" <i>Tannin</i>		368	<i>Cholera Mixture</i>	1 fl oz	250, 457
<i>Chloralamide</i>		367	" <i>Vaccine</i>		1268
<i>Chloralamidum</i>	20 to 50 gr	367	<i>Choline Distearyl Glyciophos</i>		347
<i>Chloralformamidum</i>		367	" <i>phate</i>		369
<i>Chloralose</i>		368	<i>Chondrus</i>		369
<i>Chloralum</i>		128			
" <i>formamidatum</i>		367			

CHR	Official Names in Roman, all others in Italics			
	Dose	Part		Dose
<i>Chrysa</i>		862	<i>Clark's Tiger Tails</i>	12
<i>Chrysaline</i>		860	<i>Claro de España</i>	135
<i>Christmas Poll</i>		435	<i>Clay, China</i>	700
<i>Christmas Rose</i>		506	" <i>Pocelain</i>	700
"		917	<i>Claymen's Solution</i>	104
"		917	<i>Clonal Hidratado</i>	305
<i>Chronic Acid</i>		37	<i>Cloralamido</i>	167
" <i>Anhydride</i>		37	<i>Cloratio Idiata</i>	362
<i>Chromi Trochism</i>		38	"	601
<i>Chrysosauze</i>		17	"	914
<i>Chrysosolm Crude</i>		183	" <i>Potasio</i>	914
" <i>Plaster Nails</i>		380	<i>Clorurato de Chinina</i>	901
<i>Chrysosolmum</i>	1 to 1 gr	378	" <i>Cocaina</i>	405
<i>Chrysophanic Acid</i>		784	<i>Cloruro di Ammonio</i>	144
<i>Chumbo</i>		910	" <i>de Apomonia</i>	178
<i>Chymosin</i>		549	" <i>di Calce</i>	298
<i>Chymico de Mercurio</i>		607	" <i>de Calce</i>	285
" <i>di Potasio</i>		949	" <i>Calce</i>	285
<i>Cicuta</i>		137	" <i>de Cocaina</i>	400
<i>Cicotine</i>		441	" <i>di Fide</i>	110
<i>Cigu</i>		137	" <i>Potasio</i>	521
<i>Cinco de Rhizonia</i>		580	" <i>Mercurio</i>	621
<i>Cincho</i>	1 to 5 gr	381	" <i>Mercurio al Vapor</i>	627
<i>Cinchona Calisaya</i>		129	" <i>Precipitado</i>	627
" <i>Flava</i>		382	" <i>Sublimado</i>	627
" <i>Fusca</i>		382	" <i>Monico</i>	784
" <i>Ledgeriana</i>		382	" <i>de Oro</i>	212
" <i>Officinalis</i>		382	" <i>de Oro y Sodio</i>	212
" <i>Peruviana</i>		382	" <i>Quinco</i>	901
" <i>Rubra</i>		382	" <i>Zinco</i>	1242
" <i>Succubra</i>		382	" <i>de Zin</i>	1242
<i>Cinchona Cortex</i>		382	<i>Clover</i>	1218
"		382, 1258	<i>Cloves</i>	333
"		392	" <i>Oil of</i>	334
" <i>Acidum</i>		393	" <i>Spirit</i>	107
" <i>Saboyas</i>		394	"	907
" <i>Sulphas</i>	1 to 10 gr	395	<i>Coca</i>	897
" <i>Acidus</i>		394	" <i>del Peru</i>	400
"		394	<i>Cocablatter</i>	400
" <i>Sulphas</i>	1 to 5 gr	394	<i>Cocci Folia</i>	400
" <i>Acidus</i>	1 to 10 gr	394	<i>Cocaina</i>	403
<i>Cincho quinoline</i>		394	<i>Cocaine Carbols</i>	413
(Squie)		900	" <i>Citras</i>	413
<i>Cineol</i>		499	" <i>II</i>	413
"		398	" <i>"</i>	405
"		398	" <i>Lactas</i>	413
"		398	" <i>Nitras</i>	413
"		39	" <i>Oleas</i>	413
" <i>Aldehyde</i>	1 to 2 min	344	" <i>Phenylas</i>	413
<i>Cine</i>	10 to 20 gr	399	" <i>Saboyas</i>	413
<i>Cure</i>		352	" <i>Sulphas</i>	413
" <i>Jaune</i>		349	<i>Cocainilla</i>	420
<i>Ciruelo de Espana</i>		971	<i>Cocconella</i>	420
<i>Cissampelos</i>		807	<i>Cocculus Indicus</i>	892
<i>Citarin</i>		547	<i>Coculus</i>	420
<i>Citral</i>		727	<i>Cochemille</i>	420
<i>Citras Ferruo Ammonium</i>		514	<i>Cochineil</i>	420
" <i>Ferruo Chlorus</i>		516	" <i>Solutum</i>	1309
" <i>Ferricus</i>		514	<i>Cochinilla</i>	420
" <i>cum Chinina</i>		516	<i>Cochonilla</i>	420
<i>Citrate de Fer Ammoniacal</i>		514	<i>Cocmento Antiseptico</i>	391
" <i>Magnisie desicchi</i>		752	" <i>de Quina Calisaya</i>	391
"		274	" <i>de Loja</i>	391
"		514	" <i>y Valeriana</i>	391
" <i>de Ferro Ammoniacal</i>		514	<i>Cocua Nut Swam</i>	1154
" <i>de Quina</i>		516	"	422
" <i>Latro</i>		705	"	425
" <i>Potassio</i>		947	" <i>Iodas</i>	425
<i>Citric Acid</i>		39	" <i>Phosphas</i>	424
<i>Citrine Ointment</i>		614	" <i>Saboyas</i>	425
<i>Citron</i>		722	" <i>Sulphas</i>	425
<i>Citronenschale</i>		722	<i>Codene Pastils</i>	425
<i>Citronensacure</i>		39	<i>Cod liver Oil</i>	790
<i>Citronenol</i>		722	<i>Coentro</i>	448
<i>Citronensafft</i>		727	<i>Coffena</i>	272
<i>Citrophon</i>	7 to 15 gr	875	<i>Coffeno Natrum Salicylicum</i>	277

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	Dose	Page		Dose	Page
<i>Coffeum Natri benzoicum</i>		276	<i>Confectio Sennæ</i>	60 to 120 gr	1074
" <i>Natri Salicylicum</i>		277	" <i>Sulphuris</i>	60 to 120 gr	1179
<i>Coffull's Inhalation Fluid</i>		673	" <i>Terebinthina</i>		1201
<i>Coring</i>		472	<i>Conii Folia</i>		438
<i>Cola de Pescada</i>		655	" <i>Fructus</i>		439
<i>Colantra</i>		418	<i>Conium</i>		441
<i>Colchicum</i>		432	<i>Coniæ Hydrochloridum</i>	1 to 2 gr	442
<i>Colchici Cormus</i>		426	" <i>Hydrochloridum</i>		442
" <i>Semina</i>		429	<i>Conium</i>		447
<i>Colchicina</i>	1/4 to 1 gr	432	<i>Convolvulus</i>	1 to 2 gr	443
<i>Colchicina Salicylas</i>	1/4 to 1 gr	438	<i>Convolvulus</i>		443
<i>Colchicine</i>		432	<i>Convolvulus</i>		443
<i>Colchicum</i>	2 to 5 gr	426	<i>Convolvulus</i>		700
<i>Colchique</i>		426	<i>Copahu</i>		443
<i>Colchici sal</i>		432	<i>Copallina</i>	6 to 60 min	443
<i>Cold Cream</i>	.55	1026	<i>Copaiya</i>		443
<i>Coley's Fluid</i>		1265	(<i>copaiyabalsam</i>)		443
<i>Colic Lint</i>		115	<i>Coppe</i> (See also <i>Cuprum</i>)		
<i>Colla de Pece</i>		055	" <i>Aseptol</i>		468
" <i>Piscinum</i>		055	" <i>Anticlimat</i>		910
<i>Collargol</i>		188	" and <i>Silici Albuminati</i>		405
<i>Colle de Poisson</i>		615	<i>Copulicot</i>		1017
<i>Collemplastrum Adhasivum</i>		76	<i>Coniandre</i>		448
" <i>Salicylatum</i>		76	<i>Coniandi Fructus</i>	20 to 60 gr	148
<i>Collinsonia</i>	15 to 60 gr	433	(<i>Coniandio</i>)		448
<i>Collodio</i>		976	<i>Conizuelo de Centeno</i>		483
<i>Collodion Cantaridato</i>		20	<i>Connutina</i>		491
" <i>Hæmostatic, Di Paresis</i>		977	<i>Connutina Citrus</i>	1 to 1 gr	489
<i>Collodium</i>		976	" <i>Irgotas</i>		490
" <i>Acetonum</i>		975	" <i>Hydrochloridum</i>		499
" <i>e Acido Salicylico</i>		77	<i>Corrosivi Sublimati</i>		621
" <i>Anodynum</i>		977	" <i>Dises</i>		624
" <i>Belladonna</i>		236	<i>Cortecia de Angustura</i>		469
" <i>Callosum</i>		77	<i>Cortex I rangula</i>		1011
" <i>Cantharidini</i>		321	<i>Corte a de Angostura</i>		469
" <i>Chalcium</i>		978	" <i>Granado</i>		576
" <i>Flexile</i>		976	" <i>Quilana</i>		960
" <i>Iodatum</i>		671	<i>Corallo</i>		979
" <i>Iodi</i>		671	<i>Cosil</i>		111
" <i>Iodoformi</i>		684	<i>Coscinum</i>		295
" <i>Lacto Salicylicum</i>		77	<i>Cosmoline</i>		562
" <i>Salicylicum</i>		77	<i>Coster's Paste</i>		672
" <i>Compositum</i>		77	<i>Cotarnum Hydrochloridum</i>	1 to 1 gr	845
" <i>e Zinci Chlorido</i>		77	" <i>Phthalis</i>	1/2 gr	845
" <i>Stypticum</i>		976	<i>Coto</i>		450
" <i>Tiglu</i>		461	<i>Cotoin</i>	1 to 2 gr	450
" <i>Vaseans</i>		920	<i>Coton hydrophile</i>		575
<i>Colloid Mercury</i>		606	" <i>Iode</i>		671
" <i>Silver</i>		188	<i>Cotone Absorbente</i>		575
" <i>Styptic</i>		976	" <i>Collodio</i>		975
<i>Colloidal Bismuth Oxide</i>		258	<i>Cotton</i>		575
" <i>Hæmostobin</i>		593	<i>Cotton Wool</i>		575
<i>Colloxylinum</i>		975	<i>Couch Grass</i>		1219
<i>Collyrium Alkalinum</i>		1099	<i>Couso</i>		471
" <i>Co</i>		1099	<i>Cuernicillo de Centeno</i>		485
<i>Collyre au Sulfate de Zinc</i>		1251	<i>Cramus</i>		713
<i>Collyrium Acidi Borici</i>		26	<i>Cravagem de Centeno</i>		485
" <i>et Zinci Sulphatis</i>		26	<i>Cravinho</i>		333
" <i>Opia</i>		844	<i>Cravions d Azotati d Argent</i>		186
<i>Colocynthis Pulpa</i>	2 to 8 gr	433	" <i>Mitgi</i>		186
<i>Colofonia</i>		1006	" <i>de Tannin</i>		87
<i>Colombo</i>		295	<i>Clean of Iartar, Pmited</i>		966
<i>Colophonum</i>		1006	" <i>Soluble</i>		968
<i>Coloquinte</i>		433	<i>Cremer's Bismuth</i>		247
<i>Coloquintida</i>		433	" <i>Isthargyri</i>		921
<i>Coloquintide</i>		438	" <i>Morrhue Fumicatus</i>		793
<i>Coloren de peles</i>		905	<i>Crocin (Atmann's)</i>		43
<i>Colutorio Boratado</i>		261	" <i>(Feyes)</i>		48
<i>Condurango Cortex</i>		436	" <i>(Pearson's)</i>		48
<i>Confectio Bela Recentis (Squaræ)</i>		222	<i>Cresol</i>		451
" <i>Guaiaci Composita</i>		553	<i>Cresotal</i>		455
" <i>Opia</i>		519	<i>Cresotal Carbomus</i>		455
" <i>Pipilis</i>	60 to 120 gr	901	" <i>Oleum</i>	15 to 30 gr	455
" <i>Rosa Gallie</i>		101	" <i>Isthargyri</i>	5 to 15 gr	451
" <i>Sassa</i>		101	" <i>Tannus</i>	5 to 15 gr	451
" <i>Sammum</i>	10 to 30 gr	1061	" <i>Alcalina</i>		451
			<i>Cresol</i>		451

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Dill Fruit Oil of		162	Fluorescent Typoset Salt		774
WATER		162	" Lithium Citrate		736
Diltsanum		162	" Magnesium Sulphate		754
Diluted Alcohol		1146	" Phenactin with Caffeine		875
Glucose		564	" Phenone		879
Diluting Mixture		897	" Powder of Carboid Salt		1132
Dimethylamido antipyrin		880	Sodium Citrate etc		1099
Dimethylamino Acid		1104	" Iodophate		1125
Dimethylthyl Carbonyl		158	" Sulphate		1152
Dimethyl methane diethyl sul			" Sulphate Powder		1030
phane		1172	Ibuprofen etc		124
Dioctyl ester of Paraffin		105	Iodine		847
Dimethylanthranil		1204	Irenthanthranil		91
Dioctylanthranil		975	Ika Iodophate		600
Dionine		751	Ika Iodophate		824
Diosmin		265	Ika Iodophate		323
Dioxybenzoylaminomethylacetate			Ika Iodophate		481
mine		547	Ika Iodophate		481
Di para amino phenol		417	Ika Iodophate		480
quarantine hydrochloride		1261	Ika Iodophate		480
Diphtheria Antitoxin		990	Ika Iodophate		480
Dioctylanthranil		150	Ika Iodophate		1040
Disks of Iodophate		205	Ika Iodophate		596
" " Actopine		412	" Ika Iodophate		1231
" " Cocaine		455	" Ika Iodophate		447
" " Ergotin		600	" Ika Iodophate		680
" " Homatopine		600	" Ika Iodophate		482
" " with Cocaine		653	" Ika Iodophate		124
" " Hyoscyamine		655	" Ika Iodophate		779
" " Hyoscyamine		890	" Ika Iodophate		574
" " Physostigmine		84	" Ika Iodophate		116
Disinfecting with Sulphur		1105	" Ika Iodophate		139
Dissolved Methoxybenzene		570	" Ika Iodophate		166
Dispensing Syrup		181	" Ika Iodophate		209
Distilled Water		112	" Ika Iodophate		1105
Dithion		130	" Ika Iodophate		1105
Dithion		201	" Ika Iodophate		1105
Dobell's Solution		480	" Ika Iodophate		1105
Doe amargo		195	" Ika Iodophate		1105
Donovan's Solution (Irene)		857	" Ika Iodophate		1105
Dormideura		386	" Ika Iodophate		1105
Dormol		480	" Ika Iodophate		1105
Douce amure		685	" Ika Iodophate		1105
Doves Powder		690	" Ika Iodophate		1105
" " Fluid		479	" Ika Iodophate		1105
Duboisia Myoporoides		479	" Ika Iodophate		1105
Duboisina Sulphate		479	" Ika Iodophate		1105
Duboisine		480	" Ika Iodophate		1105
Dupong Oil		480	" Ika Iodophate		1105
Dulcamara		563	" Ika Iodophate		1105
Dulcin		585	" Ika Iodophate		1105
Duotal		827	" Ika Iodophate		1105
Eucalypt Oil		529	" Ika Iodophate		1105
Faston's Pill		525	" Ika Iodophate		1105
Syrup		897	" Ika Iodophate		1105
Jau de Cannelle		258	" Ika Iodophate		1105
" " Chai		727	" Ika Iodophate		1105
" " Cologne		909	" Ika Iodophate		1105
" " Goudron		302	" Ika Iodophate		1105
" " Javelle		135	" Ika Iodophate		1105
" " Juice		82	" Ika Iodophate		1105
" " Rabel		161	" Ika Iodophate		1105
" " Destille		1026	" Ika Iodophate		1105
" " de Rose		210	" Ika Iodophate		1105
" " de Fleurs d'Orange		755	" Ika Iodophate		1105
" " Saline Iugative		755	" Ika Iodophate		1105
" " Gazeuse		312	" Ika Iodophate		1105
" " Sedative		755	" Ika Iodophate		1105
" " de Seitz		722	" Ika Iodophate		1105
Fource de Citron		578	" Ika Iodophate		1105
" " Grenadier		950	" Ika Iodophate		1105
" " Panama		275	" Ika Iodophate		1105
Effluvia Citrate		276	" Ika Iodophate		1105
" " Hy			" Ika Iodophate		1105
" " diobionide			" Ika Iodophate		1105

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<i>Extractum Lupuli</i>	5 to 11 gr	744	<i>Ferratin</i>		508
" <i>Lupulin</i>	1 to 5 gr	744	<i>Ferri Acetatis Liquor</i>	5 to 15 min	508
" <i>Malatis Ferr</i>		506	" <i>Albuminas</i>	3 to 10 gr	507
" <i>Malti</i>		268	" <i>Alginas</i>	2 to 15 gr	508
" " <i>Ferratum</i>	1 to 4 dim	270	" et Ammonii Citras	5 to 10 gr	514
" " " <i>cum Morrhu</i>		270	" " <i>Sulphas</i>	5 to 10 gr	534
" <i>Medulla Rubra</i>		761	" " <i>Tantras</i>		537
" <i>Menyanthes</i>	15 to 30 gr	773	" <i>Arsenas</i>	1 to 4 gr	508
" <i>Mezerei 1 thereum</i>		775	" <i>Arseno Citras</i>	Am	509
" <i>Nucis Vomice</i>	4 to 1 gr	815	" <i>moniaata</i>		509
" " <i>Iiquidum</i>	1 to 3 min	815	" <i>Bromidum</i>		509
" <i>Opn</i>	4 to 1 gr	835	" " <i>Cacodylas</i>	1 to 5 gr (oral) 17 min (hypodermic)	1104
" " <i>Iiquidum</i>	5 to 30 min	837	" <i>Carbonis Saccharatus</i>	10 to 20 gr	510
" <i>Papanens Iiquidum</i>	30 to 60 min	858	" <i>Chloridum</i>		521
" <i>Paren Iiquidum</i>	1 to 2 fl drim	865	" <i>Fluoridum</i>		56
" <i>Physostigmatis</i>	4 to 1 gr	887	" <i>Glycerophosphas</i>	5 to 10 gr	256
" <i>Picrotoxia Iiquidum</i>	20 to 60 min	892	" <i>Hydroxidum cum Mag</i>		15
" <i>Pini Pumiliaris</i>		901	" <i>nesu Oxido</i>		517
" <i>Piscidia</i>	1 to 5 gr	906	" <i>Hypophosphis</i>		615
" " <i>Iiquidum</i>	30 to 120 min	906	" <i>Ichthyosisulphonas</i>		519
" <i>Pomi Lcratum</i>		508	" <i>Iodidum</i>		61
" <i>Quassia</i>		979	" <i>Lactas</i>	5 to 15 gr	1136
" <i>Ahamni Frangula</i>	15 to 60 gr	1011	" <i>Meta vanadas</i>		810
" " <i>Iiquidum</i>		1011	" <i>Nucleinas</i>	5 gr	508
" " <i>Purshianil</i>		238	" <i>Peptonas</i>	5 to 10 gr	522
" " " <i>Iiquidum</i>		339	" <i>Perchloridi Iiquor Fortis</i>		521
" <i>Ihei</i>	2 to 8 gr	1014	" <i>Perchloridum</i>		526
" <i>Comp</i>		1018	" <i>Pernitratu Liquor</i>	5 to 15 min	528
" <i>Sarsa</i>		1056	" <i>Peisulphatis Liquor</i>		527
" " <i>Iiquidum</i>	2 to 4 fl drim	1056	" <i>Phosphas</i>	5 to 10 gr	530
" " <i>Compositum</i>		1056	" <i>Solubilis</i>		537
" <i>Scilla</i>		1063	" et Potassu Tartias		504
" <i>Scopola</i>		1069	" <i>Pulms</i>		530
" <i>Scalpis Cornuti</i>		457	" <i>Pyrophosphas Solubilis</i>		515
" " <i>Cornutino</i>			" et Quinin Citias	5 to 10 gr	517
" " <i>Sphacleracium</i>		491	" <i>Quinnce et Stychnine</i>		516
" " <i>Fluridum</i>		487	" <i>Citras</i>	2 to 5 gr	517
" <i>Senner Leguminosum Li</i>	1 fl drim	1070	" et Quinnce Citras Solu		516
" <i>Stiamoni</i>	4 to 1 gr	1157	" <i>biis</i>		517
" " <i>Folia</i>		1156	" <i>Stychnina Citras</i>	2 to 5 gr	530
" <i>Strophanthi</i>	4 to 1 gr	1104	" <i>Sulphas</i>	1 to 5 gr	532
" <i>Strychni</i>		816	" " <i>Exsiccatus</i>	1 to 3 gr	531
" <i>Sumbul</i>		1182	" <i>Crudus</i>		37
" <i>Supranenatum Iiquidum</i>		1184	" <i>Sulphocarbolas</i>		521
" <i>Taraxaci</i>	5 to 15 gr	1194	<i>Ferrie Chloride</i>		517
" " <i>Iiquidum</i>	1 to 2 fl dim	1194	" <i>Hypophosphate</i>		558
" <i>Tritici Iiquidum</i>	1 to 6 fl dim	1220	<i>Ferrichlor</i>		254
" <i>Triticance</i>		1230	" <i>le ruer s Snuff</i>		510
" <i>Turris Prunifolia</i>		1233	<i>Ferriol</i>		579
" " <i>Iiquidum</i>	60 to 120 min	1233	<i>Ferripyrin</i>	5 gr	537
" <i>Utrat de Cubebe</i>		464	<i>Ferrititate de Potassium</i>		503
" " <i>Fougere Muli</i>		540	<i>Ferro</i>		59
" " <i>Gaiou</i>		776	" <i>hamol</i>		537
" " <i>Iuglosse</i>		572	" <i>Kalium Tartaricum</i>		504
<i>Eye Ointments</i>		1221	" <i>Potassiato</i>		534
			" <i>Autotto dall Idrogeno</i>		590
<i>Lava de S. Ignacio</i>		655	<i>Ferrosulfat</i>		517
<i>Lava del Caldas</i>		586	" <i>terrous Hypophosphate</i>		511
<i>Lebifuge Spirit (Clutton)</i>		107	<i>lactum</i>		514
<i>Lebling's Solution</i>		108	" <i>Carbonicum Saccharatum</i>		514
<i>Leigen</i>		535	" <i>Citrum</i>		514
<i>Leibovium Pumiliatum</i>	5 to 15 gr	502	" " <i>Ammoniatum</i>		516
" <i>Bonis Pungentum</i>		505	" " <i>Chumatum</i>		514
" <i>Tarui Injessatum</i>		503	" <i>Cupulatum</i>		525
<i>Lele Maschio</i>		539	" <i>Hydro oxydatum Dulysa</i>		519
<i>Lelectinas</i>		575	" <i>tum Iiquidum</i>		519
<i>Lelech</i>		540	" <i>Iodati Saccharatum</i>		519
<i>Lelechum</i>		540	" <i>Iodatum Solutum</i>		51
<i>Leleto Crystallato</i>		27	" <i>Oxidum Saccharatum</i>	5 to 15 gr	52
<i>Leleto Doua</i>		540	" <i>Oxychloratum Solution</i>		504
<i>Lele</i>		503	" <i>ulveratum</i>		534
" <i>Leleto jati Indigeno</i>		534	" <i>hectetum</i>	1 to 5 gr	521
<i>Leleol</i>		534	" <i>Resquidatum</i>		521

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<i>Herum Sesquichloratum Cry-</i>			<i>Fluextractum Malti</i>		
<i>talatum</i>		521	" <i>Matico</i>		761
" <i>Subcarbonatum</i>		511	" <i>Mezei</i>		776
" <i>Sulphureum Præcipita-</i>			" <i>Phytolacca Radix</i>	1 to 5 min	802
<i>tum</i>		530	" <i>Podocarp</i>		697
" <i>Pututium</i>	5 to 10 gr	537	" <i>Podophylli</i>		925
<i>Ferran</i>	10 to 30 gr	508	" <i>Pruni Virginiana</i>	1 to 1 fl dr	970
<i>Festes Passifloræ</i>		858	" <i>Quassia</i>		979
<i>Feto Macho</i>		539	" <i>Quincho</i>		979
<i>Femle de Coca</i>		400	" <i>Rhamni Frangula</i>	1 to 4 fl dr	1011
<i>Femles de Belladoni</i>		224	" " <i>Purshiana Aromati-</i>		
" " <i>Bucco</i>		265	<i>cum</i>		399
<i>Fle de Calaba</i>		886	" <i>Rhex</i>	15 min	1016
" <i>de St Ignac</i>		615	" <i>Rhus</i>		1018
<i>Florolysia</i>	10-5		" <i>Loxie</i>		1023
<i>Flehi</i>		538	" <i>Salicis Arita</i>	30 to 60 min	107
<i>Fichtenharz</i>		906	" <i>Savapilla Compositum</i>		1046
<i>Ficus</i>		538	" <i>Sella</i>	1 min	1065
<i>Fenro</i>		504	" <i>Scopola</i>		1069
<i>Figos Passados</i>		535	" <i>Santalaria</i>	1 to 1 fl dr	1069
<i>Figs</i>		508	" <i>Senega</i>		1071
<i>Fique</i>		538	" <i>Senai Fructum</i>		1076
<i>Fibrene Acid</i>		510	" <i>Serpularia</i>		1078
<i>Felix Mas</i>		535	" <i>Stillingia</i>	30 min	114
<i>Filmogen</i>		977	" <i>Stannoni</i>		1156
<i>Fingerhutblatte</i>		475	" <i>Sumbul</i>		1182
<i>Finocchio</i>		340	" <i>Tryfoli</i>		1215
<i>Fish Glue</i>		655	" <i>Tritic</i>	2 fl dr	1220
<i>Meming's Tincture of Aconite</i>		94	" <i>Uta Urst</i>		1227
<i>Flexible Collodion</i>		977	" <i>Valeriana</i>	30 to 60 min	1230
<i>Flor de Cusso</i>		471	" <i>Verbati Yundis</i>		1241
<i>Flors Koso</i>		471	" <i>Verbati Opodi</i>		1244
" <i>Kusso</i>		471	" " <i>Prunifoli</i>		1244
<i>Floridena</i>		882	" <i>Verba Majoris</i>	1 to 2 fl dr	1235
<i>Flowers of Sulphur</i>		1178	" <i>Verba Santa</i>	10 to 60 min	1237
<i>Fluid Magnesia</i>		761	" <i>Fluorescent</i>		1010
<i>Fluextractum Aconiti</i>		94	" <i>Fluoriform</i>		66
" <i>Apocyn</i>	15 min	177	" <i>Fluorheum</i>		76
" <i>Arnica Radix</i>		193	" <i>Sancti Fructus</i>		610
" <i>Berberis</i>	20 to 60 min	242	" <i>Rogha di Coca</i>		400
" <i>Buchi</i>		266	" <i>Rogha di Belladonna</i>		224
" <i>Columba</i>		297	" <i>Rolia Juglandis</i>		706
" <i>Cannabis Indica</i>		317	" <i>Nutiana</i>		1190
" <i>Cassia Badreut</i>	30 to 60 min	345	" <i>Stannoni Nitrata</i>		1146
" <i>Caulophylla</i>	30 to 60 min	349	" <i>Polygoni Coca</i>		400
" <i>Composita</i>	30 to 60 min	349	" <i>Formaldehyde Ichthyol sulphur</i>		
" <i>Chilatre</i>		361	<i>ate</i>		658
" <i>Cissampel</i>	30 to 120 min	867	" <i>Formaldehydum Solutum</i>		512
" <i>Colchici Semina</i>		431	" <i>Formalin</i>		512
" <i>Collinsonia</i>	15 to 60 min	433	" <i>Formic Acid</i>		41
" <i>Convallaria</i>	8 min	443	" <i>Aldehyde</i>		512
" <i>Coto</i>	5 to 20 min	450	" <i>Formic</i>		517
" <i>Cubeba</i>	5 to 30 min	464	" <i>Formic</i>		515
" <i>Cypripedium</i>	15 min	473	" <i>Formol</i>		519
" <i>Digitalis</i>	1 to 2 min	478	" <i>Formon</i>	1 to 5 min	410
" <i>Dulcamara</i>	30 to 60 min	480	" <i>Formato Buraluo</i>		29
" <i>Ergota</i>		487	" <i>Bisulfo</i>		1115
" <i>Fru altypt</i>		498	" <i>di Calico</i>		99
" <i>Euonymus</i>		500	" <i>Di Calico</i>		21
" <i>Eupatoria</i>	20 to 60 min	501	" <i>Formoso Ferruo</i>		57
" <i>Frangula</i>	15 min	1011	" <i>de Ferro</i>		52
" <i>Gelsemi</i>	1 min	560	" <i>Sodio</i>		1113
" <i>Gentiana</i>		562	" <i>Thi Calico</i>		99
" <i>Glycyrrhize</i>		572	" <i>Porfona</i>		88
" <i>Granati</i>		575	" <i>Pothergill's Mixture</i>	oz	1066
" <i>Grindelue</i>		580	" <i>Pouge</i>		539
" <i>Guaianae</i>	30 min	585	" <i>Powles's Solution</i>		17
" <i>Hyoscyami</i>		646	" <i>Prinking</i>		1207
" <i>Jambul</i>	10 to 60 min	705	" <i>French Mixture</i>		671
" <i>Juglandis</i>		706	" <i>Preres Come's Arsenical Paste</i>		19
" <i>Kavae</i>	15 to 60 min	710	" <i>Pruri's Balsam</i>		210
" <i>Kolve</i>		712	" <i>Fructus Camali</i>		315
" <i>Krameria</i>		714	" <i>Pitidini</i>		176
" <i>Lachnanthe</i>		716	" " " "		1011
" <i>Leptandia</i>	15 min	721	" " " "		29
" <i>Lobelia</i>		741	" <i>Rhus Versiculos</i>		45
" <i>Lupulus</i>	5 to 15 min	744	" <i>Allen's Earth</i>		11

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	Do c	Page		
Glycerinum Anyli		160	Goma Tria Jacanto	16
" Atropina		304	Gonal	1044
" Belladonna		236	"	1044
" Bismuthi Carbonatis	1 to 1 fl dim	246	" Cortex	575
" Boracis		261	Gossypium	57
" Carminis		421	" Aluminis	17
" Coes		121	" Capsici	32
" Croci		459	" Depuratum	57
" Ferri Perchloridi		526	" Feratum	54
" Glycerophosphatum Compositum			" Fulminans	975
" postum		285	" Hamamelis	595
" Heronina Compositum		780	" Iodatum	671
" Hydrastis		648	" Iodoformi	664
" Hypophosphitum		291	" Krameriae	715
" Iodi		671	" Menthol	771
" Iodoformi		664	" Pungentum	57
" Ipecacuanha		689	Goudon de Houille	907
" Pancreatin		554	" vaginal	908
" Papain		836	Goulard Water	920
" Pepsini	1 to 2 fl dim	572	Gould's Extract	919
" " Fortius		973	" Lotum	920
" Plumbi Sublucetatis		920	Grasse de Lemn	100
" Resorcin		1010	Grama Franca	1220
" Soda Cinnamatis		1114	Granado	577
" Thymol Alkalinum		1200	Granati Cortex	576
" " Compositum		1209	Granatirinde	576
" "		1217	Granula Dioscoridis	19
" Bismuthi		161	Granules de Digitaline Crystal	479
" Boroglycerini		254	"	48
" Boroglycerini		25	Grasso de Cedo	1078
" Ferri Quinine et Strychninae Phosphatum	15 min	529	" Suiso	98
" Hydrastis	30 min	638	" con Benzoino	100
" Vitell		847	Greates Celandine	360
Glycerol		560	" Peruvinkle	1235
" Bismuthi Nitratis		256	Gregory's Pill	435
" ester of Guaiacol		587	" Powder	1014
" " " "		285	" Impured	1015
" " " "		291	Grenadier	577
" " " "		854	Grey Oil	60
Glycerous Hypophosphitum			" Powder	60
(Square)	1 to 2 fl dim	291	Griph's Mixture	512
" "		161	Grindelia	579
" "		160	Groundnut Oil	827
" of Ipecacuanha		689	Guaiacophol	585
" Papain		856	Guaiacotin	4 to 8 gr
" of Pepsin		873	Guaiaci Lignum	580
Glycerophosphate de Calcium		284	" Resin	5 to 15 gr
" " " "		984	Guaiacol	1 to 5 min 451, 584
Glycine		808	" Benzoate	5 to 10 gr
Glyco gelatin		556	" Cacodylate	5 gr
" gelatinum		556	" Camphorate	5 to 10 gr
Glycosal	5 to 10 gr	79	" Carbonate	5 gr
Glycyrrhizae Radix		570	" Cinnamate	5 to 10 gr
Glycyrrhizin		570	" Phosphate	5 to 10 gr
Glycyrrhizum Ammoniatum		574	" Phosphate	5 to 10 gr
Glymol		580	" Salicylate	587
Gua Powder		183	" Salol	587
Gold and its preparations See			" (Synthetic)	1 to 5 gr
also Aurum		212	" Valerianate	2 to 3 min
" Bromide		212	Guaiacool Anytol	588
" Tribromide		212	Guaiacyl	586
" Chloride of		212	Guaiacform	586
" " " "		171	Guaiacum	587
Goma Amoniac		129	Guaiamar	5 to 10 gr
" Arabica		1	Guaiaperol	904
" Guta		304	Guaiacum	587
" de Limon		482	" " "	587
Goma resina Ammoniac		129	Guaiacum	589
Gomina Ammoniac		129	"	581
" Arabica		1	"	580
" Gotta		303	"	10 to 60 gr
" Guta		304	Guaranine	587
Gomme Adraucanth		1216	Guaranne	124
" Amomaeque		129	Gum Acacia	1
" Arabique vraie		1	Gum Pastes (Unna)	4
" Gutte		303	Gummi Gutta	303

Official Names in Roman, all others in Italics

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	Dose	Page		Dose	Page
<i>Gummi Indicum</i>		588	<i>Helleboreum</i>		598
<i>Resina Gutta</i>		304	<i>Helleborum</i>		598
<i>Rubrum</i>		492	<i>Helleborus</i>		598
<i>Gummigutt</i>		303	<i>Helmistol</i>	10 to 15 gr	547
<i>Gun Cotton</i>		975	<i>Hemidesmi Radix</i>		506
<i>Gurjun Balsam</i>		213	<i>Hemlock Fruit</i>		439
<i>Gutagamba</i>		303	<i>Leaves</i>		438
<i>Gutta Percha</i>		589	<i>Hemp, Canadian</i>		177
<i>Depurata</i>		589	<i>Indian</i>		314
<i>Laminata</i>		589	<i>Henbane Leaves</i>		644
<i>Guttæ Atropinæ Sulphatis</i>		204	<i>Hepar Sulphuris</i>		832
<i>cum Cocaina</i>		204	<i>Hermophenyl</i>		607
<i>Cocainæ Hydrochloridi</i>		412	<i>Heroin</i>		778
<i>Oleosa</i>		405	<i>Hydrochloride</i>		778
<i>Creosoti (Squire)</i>		454	<i>Heronium hydrochloricum</i>		778
<i>Cupri Sulphatis</i>		468	<i>Heto Cresol</i>		1114
<i>Daturina</i>		1157	<i>Hetoform</i>		1114
<i>Euphthalmia Hydrochloridi</i>		205	<i>Hetol</i>		1113
<i>Gummi</i>		304	<i>Caffeine</i>		1114
<i>Homatropina</i>		600	<i>Hetratin</i>		547
<i>cum Cocaina</i>		600	<i>Hexahydropsura me</i>		904
<i>Hydrogenii Perovidi</i>		613	<i>Hexamethylmetramine</i>	5 to 15 gr	545
<i>Hyosina</i>		652	<i>Hexamitin</i>		700
<i>et Cocaina</i>		652	<i>Hidrastris del Canada</i>		634
<i>Phyostigmina</i>		890	<i>Hidrato de Cloral</i>		362
<i>cum Cocaina</i>		890	<i>Potasio</i>		927
<i>Pilocarpina</i>		895	<i>Sodico</i>		1096
<i>Scopolamine</i>		652	<i>Hiel de toro</i>		503
<i>Ames Chloridi</i>		1244	<i>Hiera Picra</i>		124
<i>cum Cocaina</i>		1244	<i>Hierro</i>		503
<i>Gutta</i>		304	<i>Reducido por el Hidrogeno</i>		534
<i>Gutta Test Modined</i>		1316	<i>Higos</i>		535
<i>Gutta Pills</i>		478	<i>Himrod's Cure</i>		1156
<i>Gynecol and Fincol Pancreati</i>			<i>Hinojo</i>		540
<i>(Squire)</i>		854	<i>Hypoclorito Calceo Chlorado</i>		299
<i>Gynocardia Oleum</i>		589	<i>de Calceo Empuro</i>		299
<i>Gynocardin</i>		589	<i>Sodio liquido</i>		301
			<i>Hypofosfito Calceo</i>		289
			<i>Sodico</i>		1116
<i>Haba del Calabar</i>		886	<i>Hirudines</i>		598
<i>S Ignacio</i>		658	<i>Hirudo</i>		597
<i>Hamaten</i>		592	<i>Hoffmann's Anodyne</i>		106
<i>Hamatogen</i>		593	<i>Hoja de Belladonna</i>		224
<i>Hamatoxyl Linum</i>		591	<i>Digital</i>		478
<i>Hamatoxylon</i>		592	<i>Nogal</i>		706
<i>Solution</i>		1309	<i>Taraxacon</i>		1193
<i>Hæmogallol</i>	1 to 5 gr	593	<i>Holocaine</i>		416
<i>Hæmoglobin</i>		593	<i>Hydrochloride</i>		416
<i>Colloidal</i>		593	<i>Hollow Suppositories</i>		1183
<i>Scalæ</i>		593	<i>Holunderbluthen</i>		1040
<i>Hæmol</i>	to 8 gr	593	<i>Hol kohle</i>		326
<i>Hæmostatic Colloidon</i>		977	<i>Holztheer</i>		905
<i>Hamamelidis Cortex</i>		594	<i>Homatropina</i>		600
<i>Folia</i>	1 to 5 gr	595	<i>Homatropine Hydrobromidum</i>	to 1 gr	598
<i>Hamamelis</i>		595	<i>Hydrochloridum</i>		600
<i>Hamamelis</i>		593	<i>Salicylas</i>		600
<i>Hamilton's Pill</i>		435	<i>Honey, Clarified</i>		762
<i>Hammeltalg</i>		1078	<i>Honthin</i>	10 to 30 gr	88
<i>Haid Paraffin</i>		858	<i>Hopfen</i>		742
<i>Sorp</i>		1048	<i>Hopogan</i>		848
<i>Hartsorn and Oil</i>		144	<i>Hops</i>		742
<i>Hausenblase</i>		655	<i>Hordeum Decortiatum</i>		601
<i>Hauseise</i>		1047	<i>Horseradish Root</i>		191
<i>Hausius Ammonii Chloridi</i>		146	<i>Houblon</i>		742
<i>Apomorphinæ Compositus</i>		180	<i>House Pill</i>		630
<i>Chloralamidi</i>		308	<i>Household Ammonia</i>		135
<i>Copaiba</i>		447	<i>Huile Sec Olea</i>		924
<i>Thymistrin</i>		809	<i>d Amande</i>		151
<i>Hæzeline</i>		596	<i>de Cade</i>		271
<i>Hebra. Ointment</i>		918	<i>Croton</i>		459
<i>Hedonal</i>	1 to 10 gr	1225	<i>Glyce</i>		605
<i>Hem's Pill</i>		475	<i>de Fove de Mouie</i>		790
<i>Helicho Macho</i>		539	<i>Cresotie</i>		79
<i>Helenium</i>	to 10 gr	660	<i>Phos</i>		
<i>Hellebore, White</i>		596	<i>phurée</i>		795
<i>Green</i>		1211	<i>Tusquame</i>		648
			<i>Composée</i>		648

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	Dose	Page		Dose	Page
<i>Ichland Moss</i> <i>Fucus</i>			<i>Infusum Larutimianum</i>		1076
<i>Ichthaban</i>	7 to 30 gr	658	" <i>Lupuli</i>	1 to 2 fl oz	743
<i>Ichthangan</i>		188	" <i>Matico</i>	1 to 4 fl oz	761
<i>Ichthojon</i>	10 to 20 gr	658	" <i>Pajavensis</i>		887
<i>Ichthyocolla</i>		67	" <i>Pajuna Yugmunda</i>		970
<i>Ichthyol</i>	15 to 30 gr	656	" <i>Quassia</i>	1 to 1 fl oz	978
" <i>Vismia</i>		657	" " <i>Concentratum</i>	1 to 1 fl oz	979
<i>Ichtyola</i>		655	" <i>Theri</i>	1 to 1 fl oz	1014
<i>Ichol</i>		657	" " <i>Uchabnum</i>		1014
<i>Ichast</i>		654	" " <i>Concentratum</i>		1016
<i>Igarol</i>		547	" <i>Pose Acidum</i>	1 to 1 fl oz	1022
<i>Iguatia Amara</i>		658	" " <i>Concentratum</i>		1022
<i>Ikis Iate</i>		1010	" " <i>cum lacto Nitrato</i>		1023
<i>Imperial Drink</i>		966	" <i>Scoparii</i>	1 to 2 fl oz	1066
<i>Imperial Weights and Measures</i>			" " <i>Concentratum</i>	1 to 2 fl oz	1067
<i>with Metric Equivalents</i>	xiv		" <i>Succre</i>	1 to 1 fl oz	1071
<i>Incompatibles</i> See under respective headings			" " <i>Concentratum</i>	1 to 1 fl oz	1071
" <i>Scunt</i>			" <i>Scunt</i>	1 to 1 fl oz	1071
<i>Indi Iubba</i>		322	" " <i>Compositum</i>	4 fl oz	1076
<i>Indian Gamboge</i>		305	" " <i>Concentratum</i>	1 to 1 fl oz	1076
" <i>Gua</i>		14	" " <i>cum Manna</i>		1075
" <i>Hemp</i>		548	" <i>Salinum</i>		1075
" <i>Oil of Verbena</i>		727	" <i>Scorpionum</i>	1 to 1 fl oz	1077
" <i>Senna</i>		1073	" " <i>Concentratum</i>		1075
<i>Indicators of Neutrality</i>		1509	" <i>Sinapis</i>		1085
<i>Indischer Hanf</i>		814	" <i>Linosyri</i>	1 to 1 fl oz	1216
<i>Indoform</i>	7 gr	547	" <i>Toddalia</i>	1 to 2 fl oz	1216
<i>Indurated Lead</i>		95	" <i>Urtica</i>	1 to 1 fl oz	1227
<i>Infantine</i>		710	" " <i>Concentratum</i>	1 to 1 fl oz	1227
<i>Infusa (group)</i>		659	" <i>Valeriana</i>		1230
<i>Infusion de Manis Lavante</i>		1075	" " <i>Concentratum</i>	1 to 1 fl oz	1230
" " <i>Tungate</i>		1075	" <i>Vinea Majoris</i>		1235
" <i>of Lithium</i>		714	<i>Ingluvium</i>		873
<i>de Amharbo</i>		1014	<i>Inguer</i>		1256
<i>Infuso de Rosas Composto</i>		1022	<i>Inhalations</i> See also <i>Vapores</i>		
" <i>di Sana con Manna</i>		1075	<i>Inhalatio Iodis</i>		673
<i>de Sane Composto</i>		1075	" " <i>c Conio</i>		671
<i>Intusum Abi</i>		705	<i>Injection Acid Sulphurica</i>		74
" <i>Astomata</i>	1 to 1 fl oz	124	" <i>Amylopsina (Squirt)</i>		858
" <i>Andiographulid</i>		361	" <i>Apomorphine Hypodermica</i>	5 to 10 min	180
" <i>Anthemidis</i>		180	" <i>Arsenyl</i>		1105
" " <i>Concentratum</i>	1 to 4 fl oz	169	" <i>Atoxyl</i>	{ 1 gr in 17 } { min }	1105
" <i>Armoacice Compositum</i>	1 to 2 fl oz	192	" <i>Atropina Hypodermica</i>	2 to 4 min	204
" <i>Aurantii</i>	1 to 1 fl oz	210	" " <i>et Morphina Hypo-</i>		
" " <i>Compositum</i>	1 to 1 fl oz	210	<i>dermica</i>		783
" " <i>Concentratum</i>	1 to 1 fl oz	208	" <i>Cocaine Hypodermica</i>	2 to 5 min	412
" " <i>Concentratum</i>	1 to 1 fl oz	208	" <i>Cocaine Hypodermica</i>	1 to 6 min	469
" <i>Aradachite Indica</i>	1 to 1 fl oz	210	" <i>Figat Hypodermica</i>	3 to 10 min	487
" <i>Bacchi</i>	1 to 2 fl oz	266	" <i>Ergotamine</i>		488
" " <i>Concentratum</i>	1 to 1 fl oz	296	" <i>Ferris Arsenatis</i>		509
" <i>Calumbi</i>	1 to 1 fl oz	296	" <i>Cacodylate</i>		1104
" " <i>Concentratum</i>	1 to 1 fl oz	297	" <i>Hydrargyre Biniodide</i>		612
" <i>Caryophylli</i>	1 to 1 fl oz	341	" " <i>Cyanide</i>		608
" " <i>Concentratum</i>	1 to 1 fl oz	341	" " <i>Hypodermica</i>	4 to 12 min	625
" <i>Cassia</i>	1 to 1 fl oz	342	" " <i>Iodide Iodide</i>		612
" <i>Cassia</i>	1 to 2 fl oz	349	" " <i>Solodolatus Hypo-</i>		
" <i>Chamomilla Oleum</i>		350	<i>dermica</i>	10 to 15 min	1139
" <i>Chinat</i>	1 to 1 fl oz	361	" <i>Hypocaine Hypodermica</i>	1 to 5 min	652
" " <i>Concentratum</i>		361	" <i>Ichthyol</i>		657
" <i>Chinone Acidum</i>	1 to 1 fl oz	389	" <i>Iodis</i>		671
" <i>Cocaine</i>		297	" <i>Iodoform</i>		664
" <i>Cuscuta</i>	1 to 2 fl oz	470	" <i>Morphine Hypodermica</i>	2 to 5 min	739
" " <i>Concentratum</i>		470	" <i>Morphina et Atropina</i>		
" <i>de Coiso</i>		471	<i>Hypodermica</i>	1 to 6 min	763
" <i>Digitalis</i>	2 to 4 fl oz	477	" <i>Papain Fort (Squirt)</i>		859
" " <i>Concentratum</i>	15 to 30 min	477	" <i>Prolocupina Adipatis</i>	2 to 6 min	865
" <i>Dulcamara</i>	1 to 2 fl oz	480	" <i>Quinine Hypodermica</i>		962
" <i>Eucalypti</i>	1 to 2 fl oz	487	" <i>Sol Alambroth Hypo-</i>		
" <i>Gentiana Compositum</i>	1 to 1 fl oz	562	<i>dermica</i>	10 min	626
" " " <i>Concentratum</i>			" <i>Soda Cacodylate</i>		1104
" " " <i>Concentratum</i>	1 to 1 fl oz	563	" " <i>Cinnamatis</i>	{ 16 min } { (hypodermic) }	1114
" <i>Hypocaine Oleum</i>		645	" <i>Sulphatum</i>		1251
" <i>Laborandi</i>		697	" <i>Trypsina Comp (Squirt)</i>		852
" <i>Kamille</i>	1 to 1 fl oz	714	" <i>Zinci Sulphatis</i>		1251
" " <i>Concentratum</i>	30 to 60 min	715			

Official Names in Roman, all others in Italics

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	Dose	P ₁₀₀		Dose	P ₁₀₀
<i>Sodium di Iodassio</i>		952	<i>Kino Madras</i>		711
" <i>Sodio</i>		1118	<i>Kirschschier</i>		717
<i>Juglandin</i>		706	<i>Klatschrosenblumen</i>		1017
<i>Juglans</i>		706	<i>Kobert's Decoction</i>		1067
<i>Jugo de Limones</i>		728	<i>Koch's New Tuberculin</i>		1286
" <i>Moras</i>		776	" <i>Original Tuberculin</i>		1286
<i>Juices (group) See Succa</i>		1171	<i>Koffein</i>		272
<i>Julapium Salinum</i>		948	<i>Kokum Butti</i>		552
<i>Juniper Tar Oil</i>		271	" <i>, Tre</i>		552
<i>Jumperi Oleum</i>	to min	706	<i>Kola</i>		712
<i>Jusquame noue</i>		641	" <i>Chocolate</i>		712
			" <i>Flu</i>		712
			" <i>Wajels</i>		712
			" <i>Wine</i>		712
<i>Kadcol</i>		271	<i>Kolanin</i>		712
<i>Kalabarbohu</i>		840	<i>Kollodiumwolfe</i>		975
<i>Kaladana</i>		708	<i>Kolombovri zel</i>		295
<i>Kali Canstuum</i>		928	<i>Kolophonum</i>		1006
" <i>Rusum</i>		928	<i>Koloquanten</i>		433
" <i>Solutum</i>		930	<i>Koriander</i>		448
<i>Kalidain</i>		125	<i>Koso</i>		471
<i>Kalcum Chloruum</i>		915	<i>Kosobluthen</i>		471
" <i>Situum</i>		957	<i>Kousso</i>		471
<i>Kalium See Potassium</i>		927	<i>Krameria</i>		713
" <i>Ammonium</i>		936	<i>Krimetichulix</i>	2 to 60 m	713
" <i>Bilastatum</i>		960	<i>Kraus's Catheter Lubricant</i>		35
" <i>Depuratum</i>		960	<i>Kresot</i>		450
" <i>Bromatum</i>		913	<i>Kresolum</i>		42
" <i>Carbonium</i>		943	" <i>Liquifacuum</i>		42
" <i>Depuratum</i>		943	<i>Kriotonol</i>		459
" <i>Iurum</i>		928	<i>Kryofon</i>	8 to 15 gr	876
" <i>Hydrum</i>		930	<i>Kubeben</i>		462
" <i>Solutum</i>		928	<i>Kummel</i>		331
" <i>Hydroxydatum</i>		930	<i>Kupfer sulfat</i>		466
" <i>Hydroxydatum Solutum</i>		960	<i>Kusso</i>		471
" <i>Hydroxydatum</i>		960			
" <i>Hypermanganum</i>		960	<i>La Bouillon Water</i>		19
" <i>Crystallatum</i>		960	<i>Lac Sulphuris</i>		1176
" <i>Iodatum</i>		953	<i>Lachnanthes Inctoria</i>	2 to 10 min	716
" <i>Natrium Tartarum</i>		1088	<i>Lactic Acid</i>		57
" <i>Permanganum</i>		960	<i>Lactol</i>		807
" <i>Stibio-Tartarum</i>		174	<i>Lactonaphthol</i>		807
" <i>Sulfuratum</i>		832	<i>Lactophenum</i>	5 to 15 gr	870
" <i>pro Balneo</i>		832	<i>Lactose</i>		1081
" <i>Sulfurum</i>		963	<i>Lactua</i>	5 to 15 gr	716
" <i>Tartarum</i>		964	<i>Lactuarum</i>	2 to 6 gr	716
<i>Kaliumacetat</i>		933	<i>Lact Virginat</i>		240
<i>Kaliumbromid</i>		939	<i>Lamelli Atropina</i>		203
<i>Kaliumcarbonat</i>		935	" <i>Cocaine</i>		412
<i>Kaliumchlorat</i>		944	" <i>Fuphthalmina</i>		205
<i>Kaliumcyanid</i>		940	" <i>Iyotina</i>		488
<i>Kaliumdichromat</i>		937	" <i>Homatropina</i>		800
<i>Kaliumferri stearat</i>		537	" <i>cum cocaina</i>		800
<i>Kaliumiodid</i>		952	" <i>Hyoscin</i>		653
<i>Kaliumnatriumtartrat</i>		1088	" <i>Hyoscyanina</i>		655
<i>Kaliumnitrat</i>		957	" <i>Phyostigmine</i>		890
<i>Kaliumsulfat</i>		963	" <i>Sparteine Sulphatus</i>		1066
<i>Kamala</i>		708	<i>Lana Collodii</i>		975
" <i>Depuratum</i>		708	<i>Lanoline</i>		102
<i>Kampher</i>		305	<i>Lanolina</i>		100
<i>Kaolin Massa</i>		710	<i>Lanolinum</i>		102
<i>Kaolinum</i>		709	<i>Lapides de Nitrate de Pluta</i>		186
<i>Kapose's Ointment</i>		80	<i>Lapis Divinus</i>		468
<i>Karbolaurum</i>		27	<i>Larangeira Azeda</i>		209
<i>Kardamonum</i>		328	<i>Larch Bark</i>		717
<i>Katechu</i>		346	<i>Laid</i>		98
<i>Kautschuk</i>		322	" <i>Benzoated</i>		100
<i>Kawa kawa</i>		710	" <i>Indurated</i>		98
<i>Kelene</i>		111	" <i>Oil</i>		98
<i>Kelp</i>		666	<i>Largum</i>		188
<i>Keratine</i>		710	<i>Laricus Cortex</i>		717
" <i>Solution</i>		710	<i>Larixine Acid</i>		717
<i>Kermes Mineral</i>		171	<i>Lassar's Itch Remedy</i>		806
<i>Kineurine</i>		984	" <i>Paste</i>		1248
<i>Knic Acid</i>		394	<i>Luttorio</i>		1081
<i>Knio</i>	5 to 20 gr	710	<i>Lurdano de Sydenham</i>		843
" <i>Bengal</i>		711	" <i>Vino Opinato Composto</i>		843
" <i>Eucalypti</i>	5 to 20 gr	711			

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<i>Lint, Cylin</i>		472	<i>Liquor Copasbae et Buchu et</i>		
" <i>Eucalyptus</i>		498	<i>Cubeba cum Santalo</i>		447
" <i>Iodoform</i>		664	" " <i>cum Santalo</i>		447
<i>Sathylae</i>		77	" " <i>Solubilis</i>		447
<i>Infusum Acidi Lorici</i>		26	" <i>Cosmum Concentratum</i>		297
<i>Linum</i>		79	" <i>Cresoti</i>		454
<i>Contusum</i>		70	" <i>Cresoli Saponatum</i>		40
<i>Lipase</i>		549	" <i>Cresoli Compositum</i>		43
<i>Liquefied Phenol</i>		33	<i>Glycerinatum</i>		43
<i>Sulphurous Acid Cas</i>		94	" <i>Cuscuta Concentratum</i>	1 to 1 fl drim	470
<i>Liquen (Irragan)</i>		359	" <i>Epispasticum</i>		329
<i>Islandia</i>		359	" <i>Myiobridis</i>		329
<i>Islandico</i>		359	" <i>Fragole Ammoniatum</i>	10 to 60 min	483
<i>Liquid Glucose</i>		663	" <i>Fthyl Nitritus</i>	20 to 60 min	1142
<i>Iodoform</i>		664	" <i>Luomymin et Pepsini</i>		500
<i>Liquiritia</i>		170	<i>cum Pepsino</i>		500
<i>Liquor Acidi Chromici</i>		35	" <i>Puonymini</i>	15 to 30 min	500
<i>Ardus Haller</i>		52	" <i>Herri Acetatis</i>	5 to 15 min	508
<i>Adrianae Jorvus</i>		1188	" " <i>Albuminati</i>		507
" <i>Hydrochloricus</i>		1187	" " <i>Bromidi Fortis</i>		510
<i>Ammonia</i>	10 to 20 min	132	" " <i>Chlorozandi</i>	10 to 30 min	525
" <i>Detergens</i>		175	" " <i>Dialysatus</i>	10 to 30 min	525
" <i>Jorvis</i>		181	" " <i>Hypophosphitis Ioni</i>		513
" <i>Jodati</i>		672	<i>Iodati</i>		519
<i>Ammonii Acetatis</i>	2 to 6 fl drim	143	" <i>Iodati</i>		519
" " <i>Fortis</i>		144	" " <i>Jorvis</i>		519
" " <i>Arsenatus</i>		133	" " <i>Oxychlorati</i>		525
" " <i>Arsenatus</i>		13	" " <i>Oxychloridi</i>		525
" " <i>Citratis</i>	2 to 6 fl drim	143	" " <i>Oxydati Dialysati</i>		525
" " <i>Fortior</i>		144	" " <i>Peptonati</i>		505
" " <i>(Pierlot)</i>		149	<i>cum Manganio</i>		508
" <i>Audiographidis Concentratum</i>		11	" <i>Perchloridi</i>	5 to 15 min	524
<i>Anthracidis et Papaveris</i>		163	" " <i>Fortis</i>		522
<i>Antimonii Chloridi</i>		17	" <i>Pernitritus</i>		526
" <i>Antisepticus</i>		1210	" <i>Persulphatis</i>		553
<i>Tristolochii Concentratum</i>	10 to 100 min	1078	" <i>Thiosulphatis Fortis</i>		529
" <i>Arsenialis</i>	2 to 8 min	17	<i>Subsulphatis</i>		534
" <i>Arsenici Hydrochloricus</i>	2 to 8 min	18	" <i>Itumescens</i>		1011
" <i>Arsenii Bromidi</i>		194	" <i>Bowlii</i>		17
" " <i>et Hydrargyri Iodidi</i>	5 to 20 min	195	" <i>Gutta Percha</i>		589
" <i>Atropinae Salicylatus</i>		205	" <i>Hemoglobin Co</i>		593
<i>Sulphatis</i>	to 1 min	205	" <i>Hamamelidis</i>		596
" <i>Auri et Argentii Bromatus</i>		212	" <i>Hydrogenii Peroxidi</i>	1 to 2 fl drim	640
<i>Bromadum</i>		212	" <i>Hydrargyri et Ammonii Chloridi</i>		625
" <i>Beberidis Concentratum</i>	50 to 60 min	242	" " <i>Nitratum</i>		613
" <i>Bismuthi</i>		252	" " " <i>Acidum</i>		613
<i>Concentratum</i>		254	" " <i>Perochloridi</i>	1 to 1 fl drim	624
<i>et Ammonii Citratis</i>	to 1 fl drim	252	" <i>Hypophosphitum</i>	10 to 30 min	518
" <i>Boracis</i>		261	" " <i>Compositum</i>	1 to 2 fl drim	518
" <i>Bronae Conc</i>		263	" <i>Iodi</i>		671
" " <i>Fortis</i>		263	" " <i>Carbolatus</i>		671
" <i>Bromo Chloral Compositum</i>		366	" " <i>Compositum</i>		671
" <i>Calcii Chloridi</i>	to 1 fl drim	254	" " <i>Dilutus</i>		671
" <i>Calcis</i>	1 to 4 fl oz	297	" " <i>Fortis</i>		669
" " <i>Chlorinatus</i>		300	" " <i>Glycerinus</i>		671
" " <i>Saccharatus</i>	20 to 60 min	288	" <i>Kali Arsenicosi</i>		17
" <i>Sulphuratus</i>		303	" <i>Kali Causticus</i>		930
" <i>Columbae Concentratum</i>	1 to 1 fl drim	297	" <i>Keratum</i>		710
" <i>Caoutchouc</i>		323	" <i>Krameriae Concentratum</i>	to 1 fl drim	715
" <i>Cantharidis Concentratum</i>		322	" <i>Lithii Carbonatus</i>		735
" <i>Capivi Compositum</i>		320	" <i>Magnesi Boratus</i>		28
" <i>Carbonis Detergens</i>		907	" " <i>Bromidi</i>	1 to 2 fl drim	751
" <i>Carmini</i>		421	" " <i>Carbonatis</i>	1 to 2 fl oz	751
" <i>Caulophylli et Pulsatillae</i>		349	" " <i>Citratis</i>		761
" <i>Chinita Concentratum</i>	1 to 1 fl drim	361	" <i>Meconius</i>		842
" <i>Chlori</i>		370	" <i>Morphinae Acetatis</i>	10 to 60 min	783
<i>Compositum</i>	1 fl drim	370	" " <i>Brneconatis (Squre)</i>	5 to 30 min	842
" <i>Chloroformi Compositum (Squre)</i>	5 to 10 min	376	" " <i>Hydrochloridi</i>	10 to 60 min	787
" <i>Cocci</i>		421	" " <i>Sulphatis</i>		786
" <i>Cocci</i>		421	" " <i>Nitritus</i>	10 to 60 min	790
" <i>Copasbae, Buchu et Cubeba</i>		447	" <i>Natri Caustici</i>		1086
			" <i>Natri Salicis</i>		1129
			" <i>Nitroglycerini</i>		809
			" <i>Opi Sedativus</i>	5 to 20 min	842
			" <i>Oxydi Ferrici Albuminati</i>		507

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	Dose	Page		Dose	Page
Liquor <i>Pancreaticus</i>		854	<i>Lithi Guaiacis</i>	5 gr	737
" <i>Fort (Squire)</i>	1 to 2 fl drim	853	" <i>Hypuracis</i>	5 to 15 gr	737
" <i>Pancreatis</i>		849	" <i>Tchthysulphonas</i>		656
" <i>Pectoralis</i>		874	" <i>Quinas</i>	5 to 15 gr	737
" <i>P</i>	<i>Puo</i>		" <i>Salicylas</i>	10 to 30 gr	738
" <i>P</i>		500	" <i>Theobionina Salicylas</i>	15 to 30 gr	738
" <i>Pecticus</i>		873	<i>Lithium</i>		732
" <i>Picls Carbonis</i>		907	" <i>Acidi Tartrate</i>		738
" <i>Lithanthracis</i>		907	" <i>Cafium Sulphonati</i>		738
" <i>Plumbi Subacetatis Dilu</i>		920	" <i>Dianctm</i>		738
" <i>tus</i>		919	" <i>Kinate</i>		737
" <i>" " Fortis</i>		919	" <i>Alta canadati</i>		1136
" <i>" Subacetici</i>		919	<i>Lithmus</i>		1310
" <i>" " Dulcis</i>		921	" <i>Paper</i>		1310
" <i>Potassu</i>	10 to 30 min	829	" <i>Solutum</i>		1311
" <i>" Arsenitis</i>		17	<i>Liver of Sulphur</i>		952
" <i>" (Brandish)</i>		961	<i>Iobelia</i>		740
" <i>" Chlorinata</i>		962	<i>Loblie Enfile</i>		740
" <i>" Citatis</i>		915	<i>Labelenkaut</i>		740
" <i>Potassu Arsenatis et Bio</i>			<i>Lobeline</i>		740
" <i>mulu</i>		194	<i>Lagwood</i>		591
" <i>" Citatis</i>		915	<i>Lorctm</i>		901
" <i>" Permanganatis</i>	2 to 4 fl drim	962	<i>Lacna</i>		1
" <i>Protargol</i>		150	<i>Lecophan</i>		45
" <i>Quassia Concentratu</i>	4 to 1 fl drim	975	<i>Lotio Acidi Borici</i>		26
" <i>Quinine et Strichmina</i>		528	" <i>" Carboli</i>		34
" <i>Rhei Concentratu</i>	4 to 1 fl drim	1014	" <i>" " et Boracis</i>		35
" <i>Salola Compositus</i>		1030	" <i>" Chomex</i>		39
" <i>Santal Compositus</i>		1044	" <i>" Salicylurum Boraci</i>		77
" <i>Sarsae Compositus</i>			" <i>" Tannu Sulphurata</i>		57
" <i>centiatus</i>	2 to 8 fl drim	1050	" <i>Ammonu Chloridi</i>		146
" <i>" " " " " "</i>	4 to 1 fl drim	1071	" <i>Benzoni</i>		240
" <i>Serpentina</i>	4 to 1 fl drim	1074	" <i>Bismuthi</i>		254
" <i>Sodu Chloridit</i>	4 to 2 fl drim	1078	" <i>Boracis</i>		261
" <i>Sodu Chloridit</i>	10 to 20 min	300	" <i>" cum Acido Carboli</i>		35
" <i>Sodu Arsenatis</i>	2 to 8 min	1094	" <i>Calamina</i>		280
" <i>" Boratis Compositus</i>		281	" <i>Calu Sulphurati</i>		303
" <i>" Carbolatis</i>		35	" <i>Crimals</i>		135
" <i>" Ethylatis</i>		1116	" <i>Hydrargyri Acetica</i>		625
" <i>" Hydrozatis</i>		1087	" <i>" Flava</i>		624
" <i>" Phosphatis Compositus</i>		1126	" <i>" Nigri</i>		629
" <i>" " " " " "</i>		1134	" <i>" Pochloridi</i>		625
" <i>Styck</i>			" <i>" " Acidi</i>		625
" <i>" " " " " "</i>	2 to 8 min	1168	" <i>Pancreatica Fort (Squire)</i>		854
" <i>Sublimati Corrosivi (Van</i>			" <i>Papain</i>		866
" <i>Stieten)</i>		624	" <i>Plumbi</i>		921
" <i>Suparenalis Hamostati</i>			" <i>" Acetatis</i>		913
" <i>cus</i>		1185	" <i>" Evaporatis</i>		921
" <i>Taraxaci</i>		1195	" <i>" " cum Morphu</i>		921
" <i>" " " " " "</i>		1210	" <i>" Lactatis</i>		922
" <i>" " " " " "</i>		1210	" <i>" " cum Opio</i>		921
" <i>Thyroides</i>		1213	" <i>" " et Picis</i>		907
" <i>" " " " " "</i>		1214	" <i>" " Sulphuris</i>		1177
" <i>" " " " " "</i>	30 to 60 min	1216	" <i>Resorcinu Compositu</i>		1010
" <i>Toddalce</i>	30 to 60 min	1216	" <i>Rubia</i>		1251
" <i>Tolutanus pro Syrupu</i>		220	" <i>Sodu Hyposulphitis</i>		1154
" <i>Tumtium</i>	4 to 2 min	800	" <i>Staphysagru</i>		1153
" <i>Trypsina Comp (Squire)</i>	1 to 2 fl drim	863	" <i>Sulphuris</i>		1177
" <i>" " " " " "</i>		144	" <i>" " " "</i>		851
" <i>" " " " " "</i>		1243	" <i>" " " "</i>		1240
" <i>" " " " " "</i>		106	" <i>" Chloridi</i>		1241
" <i>" " " " " "</i>		572	" <i>" Ozidi</i>		280
" <i>" Root</i>		570	" <i>" Sulphatis</i>		1261
" <i>Lino de los Valles</i>		443	<i>Lotio u l Acetate de Plomb</i>		921
" <i>Lintern</i>		28	" <i>" Ammoniacale Camphre</i>		312
" <i>Litargyro</i>		910	" <i>" " " "</i>		718
" <i>Litharge</i>		916	" <i>" " " "</i>		1103
" <i>Lithu Water</i>		735	" <i>Lozenges Cream</i>		638
" <i>Lithi Benzozas</i>	15 to 30 gr	737	" <i>Lozenges See Trochisci</i>		1220
" <i>" Butyris</i>	5 to 15 gr	738	" <i>" Bases for</i>		1220
" <i>" Bromidum</i>	5 to 15 gr	737	" <i>" " " " "</i>		1220
" <i>" Citonas</i>	2 to 5 gr	732	" <i>" " " " "</i>		671
" <i>" Citras</i>	5 to 10 gr	735	" <i>" " " " "</i>		671
" <i>" " Effervescens</i>	60 to 120 gr	736	" <i>" " " " "</i>		144
" <i>" " Lavatunus Efferves</i>		737	" <i>" " " " "</i>		35
" <i>" cens</i>			" <i>" " " " "</i>	2 to 5 gr	743
			" <i>" " " " "</i>		742

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	Dose	Page		Dose	Page
<i>Lupulus</i>		742	<i>Manganese Sulphas</i>	1 to 5 gr	768
<i>Lut. & Ointment</i>		612	<i>Mangan. Dioxidum Praecipitatum</i>		757
<i>Lycetol</i>	5 to 10 gr	905	<i>Manganum Hyperoxidatum</i>	60 gr to 1 oz	757
<i>Lycopodium</i>		744	<i>Manna</i>		759
<i>Lysidine</i>	5 to 60 min	905	<i>Depurata</i>		759
<i>Acid Tartrate</i>	1 to 30 gr	905	<i>Mannite</i>		759
<i>Lysosform</i>		547	<i>Manitol Hexanitate</i>	1 to 2 fl dim	780
<i>Lysol</i>		14	<i>Manteca de Cacao</i>		120
<i>Lythargyrum</i>		917	<i>Cerdo</i>		98
			<i>O Aceite concreto de Vuc.</i>		800
<i>Mad dog Skullcap</i>		1099	<i>Moscada</i>		167
<i>Magma Magnesii</i>	1 fl dim	755	<i>Manilla ordinaria</i>		187
<i>Magnesia Fluid</i>		751	<i>Romana</i>		760
	5 to 30 gr		<i>Masanta</i>		215
	(repeated)		<i>Margosa Bark</i>		295
	5 to 60 gr	74	<i>Mari gold Common</i>		815
	(single)		<i>Marijuana</i>		172
		75	<i>Marmelo</i>		123
	5 to 30 gr		<i>Marshall Hall's Pill</i>		124
	(repeated)		<i>Marsmallow Root</i>		513
	5 to 60 gr	747	<i>Massa Ferri Carbonatis</i>		601
	(single)		<i>Hydriopyi</i>		710
		748	<i>Kaolin</i>		885
<i>Usta Ponderosa</i>		747	<i>Paraffinum</i>		780
<i>Magnesia Calcini & pesante</i>	5 to 15 gr	755	<i>Mastix and Chloroform</i>		760
<i>Magnesium Benzoas</i>	15 to 30 gr	26	<i>Mastic</i>		780
<i>Boro Citras</i>	1 to 2 fl dim	751	<i>Matico</i>	50 to 120 gr	167
<i>Bromidi Liquor</i>		1104	<i>Matricaria</i>		1317
<i>Cnidias</i>	5 to 30 gr		<i>Mauveri & Test</i>		211
	(repeated)		<i>Measures Metric</i>		21
	5 to 60 gr	748	<i>and Weights of British</i>		775
	(single)		<i>Pharmacopœia</i>		842
	5 to 30 gr		<i>Mercurion</i>	1 to 1 gr	1048
	(repeated)		<i>Mercurii Periodolum</i>		761
	5 to 60 gr	750	<i>Methiurische Seife</i>		761
	(single)		<i>Midulla Lœvis depurata</i>		191
<i>Citrius Liquor</i>	5 to 10 fl oz	751	<i>Rubia</i>		1082
<i>Citrus Potio</i>		752	<i>Medullary Glyceride</i>		644
<i>Icthyosulphonas</i>		655	<i>Meerretig</i>		261
<i>Salicylas</i>	50 to 100 gr	756	<i>Meerzwabel</i>		769
	50 to 120 gr		<i>Mermendro</i>		771
	(repeated)		<i>Mel Potacis</i>		772
	4 to 1 oz	752	<i>Depuratum</i>		465
	(single)		<i>Rosa</i>		782
	60 to 240 gr		<i>Mellite Cumeni</i>		782
	(repeated)		<i>Mellito Simplex</i>		576
	1 to 1 oz	754	<i>Melogramato</i>		464
	(single)		<i>Melon Pumpkin Seeds</i>		472
<i>Sulphas</i>	50 to 50 gr	756	<i>Menbrillo</i>		768
<i>Sulphocarbolic</i>		37	<i>Menthae Crispæ Oleum</i>	to 5 min	764
<i>Magnesium</i>		745	<i>Menthae Piperitæ Oleum</i>	to 5 min	764
<i>Citricum Effervescentis</i>	1 to 3 gr	752	<i>Viridis Oleum</i>		771
<i>Gynocadate</i>		690	<i>Menthene</i>		771
<i>Lactate</i>		752	<i>Menthol</i>		772
<i>Oxide Light</i>		745	<i>Plaster</i>		772
<i>Heavy</i>		717	<i>Snuff</i>		772
<i>Mari & Pills</i>		123	<i>Valerianate</i>	5 to 10 min	772
<i>Malakin</i>	10 to 20 gr	576	<i>Mentholate</i>		772
<i>Male Fern</i>		538	<i>Mentholum</i>		770
<i>Mallera</i>		1268	<i>Menthosol</i>	6, 772	
<i>Malt</i>		268	<i>Menthosol</i>		773
<i>Extract</i>		288	<i>Mentol</i>		760
<i>Liquid</i>		270	<i>Mentolo</i>		769
<i>with Cod Liver Oil</i>		270	<i>Menyanthes</i>	to 6 fl oz	773
<i>Liquid</i>		270	<i>Mercuroamine</i>		606
<i>Manna</i>		750	<i>Mercure Purpur</i>		601
<i>Mandelol</i>		151	<i>Mercurial Cream</i>		604
<i>Mandole Amare</i>		150	<i>Lotion Black</i>		629
<i>Doli</i>		154	<i>Yellow</i>		624
<i>Manganese</i>		757	<i>Plaster</i>		603
<i>Manganum et Ferri Citras</i>	1 to 5 gr	758	<i>Muriatic Ammonium Chloride</i>		621
<i>Hypophosphis</i>		757	<i>Chloride</i>		610
<i>Oxulum Depuratum</i>	10 to 30 gr	756	<i>Iodide</i>		616
<i>Phosphas</i>		757	<i>Oxide</i>		619
<i>et Sodii Citras</i>	1 to 5 gr	756	<i>Oxide, Red</i>		619

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	Dose	Page		Page
Mitum & Phelus		540	Mochus	5 to 10 gr
" Gentiana	1 to 1 fl oz	563	" <i>Isucatus</i>	795
" " Iseda		563	Moss, Iceland	796
" " Alkalina		563	" " Irish	799
" " cum Soda		563	Mortaric	1080
" <i>Glycerhiza Composita</i>	2 fl drim	574	Mortaza	1079
" Guaiac	1 to 1 fl oz	583	Mountain Danson	1079
" <i>Homatophis cum Catichis</i>		348	Moutarde	1079
" <i>Ipecacuanha Amoniacata</i>		690	Mouth and Nose Protector (for poisonous and injurious trades)	875
" " <i>Sativa</i>		690	Mucilage of Moss	369
" " cum Soda		690	" " <i>Starch</i>	101
" <i>Jalap cum I heo</i>		704	Mucilago Acreta	1 to 4 fl drim
" <i>Jatropha</i>		340	" " <i>Cydonia</i>	472
" <i>Magnesia e Phos</i>		751	" " <i>Salep</i>	1035
" <i>Melilotaceantidis</i>		502	" " <i>Pasacanthus</i>	1218
" Moschi	1 to 1 fl oz	796	" " <i>Umbi</i>	1221
" <i>Mucilaginoso</i>		4	Mudar	290
" <i>Olea Ricini</i>	1 to 1 fl oz	1020	Mughetto	443
" " <i>Santali</i>		1044	Muguet	440
" <i>Paraldehyde</i>		867	Mulberry Juice	776
" <i>Peppum Composita</i>		873	Muratic Lithi	110
" " cum <i>Fritrato Malti</i>		873	Muse	795
" <i>Potassii Chloratis</i>		947	Muscade des Moluques	797
" " <i>Citrus Pfefferis</i>		949	Muschio	796
" " <i>Iodidi</i>		957	Musk	795
" " " <i>Allalina</i>		957	Muscatum	797
" " " <i>et Stramoni</i>		957	Mustard	1079
" <i>Quinnia</i>		1006	" " Oil of Volatile	1082
" " cum <i>Puro</i>		1006	" " <i>Papiri</i>	1082
" <i>Juhi cum Soda</i>		1016	Muttektion	438
" " <i>et Soda</i>		1016	Mydnasne	204
" <i>Sativa</i>		959	Mydriac	433
" " <i>Anodyni</i>		959	Nylidius	317
" " <i>Laxans</i>		757	<i>Myrsichis Ilex of 1 drim</i>	5 to 10 min
" <i>Santali Composita</i>		1044	Myristic	5 to 15 gr
" " cum <i>Morphine</i>		1011	<i>Myristic Adept</i>	809
" <i>Sammone</i>		1001	Myrobalanum	5 to 60 gr
" <i>Sidit</i>		1006	<i>Myrono Acid</i>	1079
" " <i>Composita</i>		1065	Myrron	1079
" " <i>et Ipecacuanha</i>		1065	Myrrha	800
" " " <i>Opi</i>		1065		
" <i>Sonne Composita</i>	1 to 2 fl oz	1074	<i>Nafatalan</i>	804
" <i>Sodae Composita</i>		1100	<i>Nafatalolo</i>	804
" <i>Spiritus Vini Gallici</i>	1 to 2 fl oz	1152	<i>Naphthalene</i>	803
" <i>Stychnine Acida</i>		1169	<i>Naphthum Petrachloritum</i>	2 to 10 gr
" <i>Sulphurea Acida</i>		82	<i>Naphthalinum</i>	2 to 5 gr
" <i>Tussis Rubia</i>		378	" <i>Præcipitatum</i>	803
" " <i>Concentrata</i>		378	<i>Naphthalol</i>	2 to 8 gr
Mitigatio Cautic		186	Naphthol	5 to 10 gr
Mortui for Cholera	20, 456		" <i>Camphor</i>	807
Moelle de Boeuf purpure		761	<i>Nasania Anas</i>	206
Mollin		1034	<i>Nasena</i>	to 2 gr
Monobromatid Camphor		313	<i>Nasotile</i>	111
Monobromkamphur		313	<i>Nasotina</i>	844
Mono ethyl Morphine Hydrochloride	1 to 5 gr	781	Narcot	1 to 1 gr
<i>Monosulphure Acid Glycerin</i>		79	<i>Nasol</i>	810
Monsel's Salt		534	<i>Natadon</i>	120
" <i>Solution</i>		534	<i>Natrio Kalium Tartaricum</i>	1088
Morfinum		776	<i>Natrum</i> See Sodium	10
Mors Succus		776	" <i>Causticum</i>	71
Morphina	1 to 1 gr	776	" <i>Hydricum</i>	
Morphina Acidis	1 to 1 gr	782	" <i>Solutum</i>	
" <i>Hydrobromatum</i>	1 to 1 gr	784	" <i>Hydroxydatum</i>	
" <i>Hydrochloratum</i>	1 to 1 gr	784	" <i>Solutum</i>	
" <i>Lactis</i>	1 to 1 gr	785	" <i>Hypochlorosum Solutum</i>	
" <i>Sulphas</i>	1 to 1 gr	785	" <i>Sulphurum Crystallatum</i>	
" <i>Tutias</i>	1 to 1 gr	785	" <i>Depuratum</i>	
Morphine Scopolanine Anas		650	" <i>Succum</i>	
<i>Morphinum Duetetylum</i>		778	" <i>Sulpho Ichthyolium</i>	
" " <i>Hydrochloricum</i>		779	" <i>Sulphurum</i>	
Morhuu & Olcum	1 to 4 fl drim	790	" <i>Succum</i>	
Morrhuel	3 gr	795	<i>Natrumacetat</i>	
Morton's Fluid		671	<i>Natrumarsenat</i>	
			<i>Natrumbenzoat</i>	

NAT		Official Names in Roman, all others in Italics.	
	Dose	P.L.C.	
<i>Natriumbicarbonat</i>		1096	<i>Nosophin</i> 5 gr 882
<i>Natriumboracat</i>		2 40	<i>Nosospirin</i> 79
<i>Natriumbromid</i>		1100	<i>Novaldine</i> 415
<i>Natriumbicarbonat</i>		1107	<i>Noz Moschada</i> 798
<i>Natriumchlorid</i>		1109	" <i>Yomua</i> 811
<i>Natriumhypophosphit</i>		1116	<i>Nuclein</i> 809
<i>Natriumiodid</i>		1118	<i>Nuclein Acid</i> 809
<i>Natriumphosphat</i>		1123	<i>Nucleol</i> 809
<i>Natriumsulfat</i>		1126	<i>Nuc Moscada</i> 797
<i>Natriumsulfat</i>		1130	<i>Nursery Hair Lotion</i> 113
<i>Nebula Acid Tannic</i>		87	<i>Nutric</i> 797
" <i>Adrenalin (Squint)</i>		1157	" <i>Oil of</i> 799
" <i>Adrenin</i>		1187	<i>Nut Moschada</i> 798
" <i>cum Cocaina</i>		1185	" <i>Yomua</i> 1 to 4 gr 810
" <i>Alkalina</i>		1100	
"		1100	<i>Oak Bark</i> 979
"		261	<i>Olivares Quicksilver</i> 616
"		412	<i>Ointments See Unguenta</i> 1221
" <i>Composita</i>		405	<i>Ole (group)</i> 873
" <i>Oleosa</i>		405	<i>Oleat de Moruro</i> 616
" <i>Eucalypt</i>		498	<i>Oleat of Immunia</i> 135
" <i>et Menthol et Cocaina</i>		405	<i>Oleat</i> 824
" <i>Pain</i>		405	<i>Oleatum Aconitina</i> 97
" <i>Estacti Suprasenalis</i>		1189	" <i>Atropina</i> 901
" <i>Guaiacol et Menthol</i>		585	" <i>Cocaine</i> 405
" <i>Iodi Composita</i>		671	" <i>Quinnia</i> 982
" <i>et Menthol</i>		671	" <i>Veratrine</i> 1243
" <i>Iodoform</i>		664	<i>Oleic Acid</i> 61
" <i>Menthol</i>		772	<i>Oleum Aconitina</i> 97
" <i>cum Cocaina</i>		772	" <i>Atropina</i> 901
" <i>Menthol Composita</i>		772	" <i>Hydargyro</i> 616
" <i>Sodi Chloridi Composita</i>		1112	" <i>Quinnia</i> 982
" <i>Suprasenalis et Cocaine</i>		1157	" <i>Veratrine</i> 1243
" <i>in Aqua</i>		222	<i>Oleusare</i> 60
" <i>in Aqua</i>		213	<i>Oleo See Olea</i> 824
<i>Nelkenöl</i>		834	" <i>de Bacchari</i> 791
<i>Nervocidine</i>		417	" <i>Calao</i> 1203
	1 to 5 gr	8	" 791
	1 to 1 gr	807	" 720
<i>Nickel</i>		808	" <i>Noz Moschada</i> 800
" <i>Carbonyl</i>		807	<i>Oleocroton</i> 455
<i>Nicotine</i>		808	<i>Oleocroton Aspidu</i> 540
" <i>Salicylate</i>		1190	" <i>Capivi</i> 1 to 1 min 326
<i>Nirvanin</i>		417	" <i>de Copaiba</i> 414
<i>Nitras Argenticus Mitigatus</i>		186	" <i>Cubibe</i> 5 to 30 min 464
" <i>Kalium</i>		958	" <i>Lupulini</i> 1 to 5 gr 744
<i>Nitro Argentico Cristallizado</i>		184	" <i>Piperis</i> 1 gr 905
" <i>Mitigado</i>		186	" <i>Zingiberis</i> 1257
" <i>de Argento</i>		184	<i>Oleosacchari</i> 824
" <i>de Argento Fuso con</i>			<i>Oleum Adepsini</i> 860
" <i>Nitro de Potassio</i>		186	" <i>Amygdalo</i> 1 to 4 fl drim 150
" <i>(sub) Benzoato</i>		250	" <i>Aman Essentiali</i> 162
" <i>Mercurio</i>		613	" <i>Isent Persu</i> 152
" <i>Acido</i>		613	" <i>Anethi</i> 1 to 2 min 162
" <i>Liquido</i>		613	" <i>Anisi</i> 1 to 3 min 164
" <i>di Potassio</i>		957	" <i>Anthemidis</i> 1 to 3 min 168
" <i>Potassio</i>		957	" <i>Anthos</i> 1026
<i>Nitre</i>		957	" <i>Arachis</i> 527
"		61	" <i>Asepticum</i> 82
"		64	" <i>Aurantii Corticis</i> 208
<i>Nitras Athylus cum Spiritu</i>		1140	" <i>Flor an</i> 211
<i>Nitrate of Amyl</i>		155	" <i>Betula</i> 242
<i>Nitrate of Amile</i>		155	" <i>Lenta</i> 553
<i>Nitroglycerin</i>	1 to 1 gr	808	" <i>Betulinum Rectificatum</i> 3 to 8 gr 242
" <i>Tablets</i>		809	" <i>Cacao</i> 120
<i>Nitro Hydrochloric Acid, Di</i>			" <i>Cadmum</i> 271
" <i>luted</i>		64	" <i>Cajupati</i> 1 to 3 min 278
<i>Noce Moscata</i>		797	" <i>Camphore Essentiali</i> 342
" <i>Yomua</i>		810	" <i>Camphoratum</i> 308
<i>Noce de Galla</i>		550	" <i>Forde</i> 308
<i>Noce Vomique</i>		810	" <i>Cantharidatum</i> 321
" <i>Acid</i>		79	" <i>Cardamomi</i> 330
" <i>Acid</i>		1112	" <i>Carm</i> 1 to 3 min 332
<i>Normal Serum for Alimenta</i>		1205	" <i>Caryophylli</i> 1 to 3 min 334
<i>tion</i>		158	" <i>Cassia</i> 1 to 3 min 343
<i>Norvarian</i>			" <i>Chamomille</i> <i>Camphora</i> 168

Official Names in Roman, all others in Italics

OXY

	Dose	Price		Dose	Price
<i>Decum Chamomilla Infusum</i>		168	<i>Olum Ligni</i>		460
" <i>Chaulmoogra</i>		590	" <i>Induratum</i>	2 to 5 min	1229
" <i>Chloroformi</i>		575	<i>Oleo</i> See <i>Olea</i>		824
" <i>Cinereum</i>	1 to 2 min	603	" <i>Cadino</i>		271
" <i>Cinnamomi</i>	1 to 3 min	597	" <i>di Crotontiglio</i>		459
" <i>Citri</i>		723	" <i>Regato di Merli eo</i>		790
" <i>Copulati</i>	5 to 20 min	446	" <i>Mandole Dolci</i>		151
" <i>Coumudi</i>	1 to 3 min	149	" <i>Olive</i>		824
" <i>Crotonis</i>	1 to 1 min	459	" <i>Iticno</i>		1018
" <i>Tructum</i>		400	<i>Olive Oil</i>	1 to 1 fl oz	824
" <i>Cubeli</i>	5 to 20 min	463	" <i>Stalised</i>		425
" <i>Declina</i>		860	<i>Ohrenni</i>		1824
" <i>Frugantis Canadensis</i>	5 to 10 min	191	<i>Olivari Contar</i>		1049
" <i>Lacrypli</i>	1 to 3 min	194	<i>Ohno</i>		231
" <i>Maculati vari Citri</i>			<i>Onquent Aggyptiac</i>		465
" <i>odora</i>		419	<i>Opi Punctura cum Acido Ben</i>		310
" <i>Tranquila</i>	5 to 15 min	541	" <i>zouo</i>		827
" <i>Tranquilla</i>	5 to 15 min	552	<i>Opiu</i>	1 to 2 gr	827
" <i>Tranquilla Citri</i>		727	<i>Opium</i>		827
" <i>Gymnadi</i>		549	" <i>di Smyru</i>		841
" <i>Homatipapiri e Conaria</i>		600	" <i>Group of Preparations</i>		1053
" <i>Hyoscyami</i>		647	" <i>with Proportions</i>		946
" <i>Compositum</i>		648	<i>Opodilol</i>		956
" <i>Johorini coctum</i>		648	" <i>Iodatum</i>		1053
" <i>Infusum</i>		648	" <i>Liquidum</i>		827
" <i>Jecoris</i>		791	<i>Opino</i>		1287
" <i>Aselli</i>		790	<i>Opionans</i>		211
" <i>cum Iodo Lei</i>		791	<i>Opionans</i>		210
" <i>feratum</i>		790	" <i>Pcel, Bitter</i>		208
" <i>Iodatum</i>		791	" <i>Wine</i>		207
" <i>Juniperi</i>	1 to 3 min	706	<i>Orchitic Fluid, Lion n Se</i>		1139
" <i>Empyricum iticum</i>		371	" <i>guard</i>	1 to 5 gr	846
" <i>Empyricum iticum</i>		715	" <i>Hydrochloride</i>	2 to 8 gr	846
" <i>Empyricum iticum</i>	1 to 3 min	719	" <i>Annali</i>	5 to 10 gr	846
" <i>Empyricum iticum</i>	1 to 3 min	722	<i>Orge Perl</i>		801
" <i>Empyricum iticum</i>	1 to 3 min	730	<i>Oro u</i>		871
" <i>Empyricum iticum</i>		731	<i>Orphol</i>		258
" <i>Empyricum iticum</i>		731	<i>Orthochoxy ana Monobenzoyle</i>		161
" <i>Empyricum iticum</i>		731	" <i>amidochinolun</i>		564
" <i>Empyricum iticum</i>		731	<i>Orthobenzoic sulphide</i>		416
" <i>Empyricum iticum</i>		731	<i>Orthoform</i>		415
" <i>Empyricum iticum</i>		731	" <i>Hydrochloride</i>	1 to 5 gr	415
" <i>Empyricum iticum</i>		731	" <i>New</i>		415
" <i>Empyricum iticum</i>		731	<i>Osmic Acid</i>		67
" <i>Empyricum iticum</i>		731	<i>Osmiumsulphide</i>		298
" <i>Empyricum iticum</i>		731	<i>Ossido di Calcio</i>		67
" <i>Empyricum iticum</i>		731	" <i>Magnesio</i>		745
" <i>Empyricum iticum</i>		731	" <i>Mercurio Gallo</i>		617
" <i>Empyricum iticum</i>		731	" <i>Rosso</i>		619
" <i>Empyricum iticum</i>		731	" <i>di Zinco</i>		1245
" <i>Empyricum iticum</i>		731	<i>Ossigeno</i>		848
" <i>Empyricum iticum</i>		731	<i>Ostindischer En ran</i>		360
" <i>Empyricum iticum</i>		731	<i>Otto of Rose</i>		1023
" <i>Empyricum iticum</i>		731	<i>Orubani</i>		1165
" <i>Empyricum iticum</i>		731	<i>Ourouparia Gambir</i>		345
" <i>Empyricum iticum</i>		731	<i>Ova Albumen</i>		846
" <i>Empyricum iticum</i>		731	" <i>Vitellus</i>		847
" <i>Empyricum iticum</i>		731	<i>Ox Bile, Purified</i>		502
" <i>Empyricum iticum</i>		731	<i>Oxalate de Cero</i>		354
" <i>Empyricum iticum</i>		731	<i>Oxalic Acid</i>		67
" <i>Empyricum iticum</i>		731	<i>Oxalium e</i>		68
" <i>Empyricum iticum</i>		731	<i>Oxaphor</i>		313
" <i>Empyricum iticum</i>		731	<i>Oxithylpara Acetanilide</i>		873
" <i>Empyricum iticum</i>		731	<i>Oxido Antimonoso Precipi</i>		169
" <i>Empyricum iticum</i>		731	" <i>tado</i>		525
" <i>Empyricum iticum</i>		731	" <i>de Ferro Dialysado</i>		745
" <i>Empyricum iticum</i>		731	" <i>Magnesio</i>		619
" <i>Empyricum iticum</i>		731	" <i>Mercurio</i>		617
" <i>Empyricum iticum</i>		731	" <i>Amarillo</i>		619
" <i>Empyricum iticum</i>		731	" <i>Rodo</i>		917
" <i>Empyricum iticum</i>		731	" <i>de Plomo</i>		925
" <i>Empyricum iticum</i>		731	" <i>Potasio</i>		1245
" <i>Empyricum iticum</i>		731	" <i>Zinco</i>		241
" <i>Empyricum iticum</i>		731	<i>Oxyacanthine</i>		

OXY

Official Names in Roman, all others in Italic

	Doct.	Pt.		Doct.	Pt.
<i>Oxyampbon</i>	15 to 30 gr	41	<i>Tanaphenol</i>	1 to 5 grains	876
<i>Oxychinasceptol</i>		991	" <i>Aquatic</i>		876
<i>Oxyde d'Argent</i>		190	" <i>Amorphate</i>		875
" <i>de Calcium</i>		298	" <i>Camphorate</i>		876
" <i>Magnesium</i>		745	" <i>Citrate</i>		875
" <i>Mercuri Tannic</i>		617	" <i>Lactate</i>		876
" <i>" Rouge</i>		619	" <i>Methyl Glycollate</i>		876
" <i>Plomb Fondue</i>		916	" <i>Sulphate</i>		876
" <i>" Zinc</i>		1215	" <i>Quinine ethyl Carbonate</i>		876
<i>Oxyde de Chumbo</i>		917	" <i>Vanillin ethyl Carbonate</i>		876
" <i>Mercurio</i>		619	<i>Paragoric</i>		309
<i>Oxydum Hydragryni Plurimum</i>		617	" <i>Elisir</i>		309
" <i>" Rubrum</i>		619	" <i>Stoch</i>		311
" <i>Hydragryriacum</i>		619	<i>Paracitradix</i>		307
" <i>" Flavum</i>		617	<i>Paranol</i>		303
" <i>" Precipitatum</i>		617	" <i>Isoscar</i>		30
" <i>" Pondicosum</i>		745	" <i>Liquid</i>		30
" <i>" Semivivum</i>		917	" <i>Spirituale</i>		30
" <i>Stibium</i>		109	" <i>Wool Fat</i>		30
<i>Oxygen</i>		547	<i>Paragen Guaiacolis</i>		353
<i>Oxyhmoglobin</i>		593	" <i>Ichthamolis</i>		67
<i>Oxymel</i>	1 to 2 fl drim	763	" <i>Iodi</i>		673
" <i>Ipecacuanha</i>		690	" <i>Liquid</i>		864
" <i>Scilla</i>	1 to 1 fl drim	1064	" <i>Sulphatatum</i>		78
" <i>Urgine</i>		1064	<i>Parogenum</i>		304
" <i>de Verdete</i>		465	" <i>Chloroformi Camphora</i>		378
" <i>" "</i>		1046	" <i>tum</i>		378
" <i>" "</i>		1063	" <i>Creosoti</i>		455
<i>Oxypteris Hydrochloric</i>		1068	" <i>" "</i>		271
" <i>Sulphate</i>		1068	" <i>" "</i>		498
<i>Ozone</i>		848	" <i>" "</i>		606
<i>Ozone Ether</i>		641	" <i>" "</i>		673
			" <i>Iodoformi</i>		605
<i>Palo de Campeche</i>		591	" <i>" Diiodatum</i>		607
<i>Panama Bark</i>		980	" <i>Mentholis</i>		778
<i>Pancreas Enzymes</i>		848	" <i>Naphtholis</i>		800
<i>Pancreatic Diastase</i>		851	" <i>Picea</i>		916
" <i>Emulso</i>	1 to 4 fl drim	855	" <i>Synsyrin</i>		864
<i>Pancreatic Solution</i>		849	" <i>Sulphuris</i>		1150
<i>Pancreatin</i>		549	" <i>" Compositum</i>		1150
<i>Pancreatina</i>		550	" <i>Terebinthina</i>		717
" <i>Medicinalis</i>		850	<i>Paroleine</i>		500
<i>Pancreatinum</i>		549	<i>Parish's Chemical Food</i>		549
<i>Pancreatised Fat</i>	1 to 4 fl drim	855	<i>Pasta Acidi Salicylici</i>		77
<i>Pankreatinum</i>		550	" <i>Copasbi</i>		148
<i>Papain</i>	2 to 10 gr	555	" <i>Hamamelidis</i>		596
<i>Papaverina</i>	1 to 1 gr	545	" <i>Ichthamolis</i>		67
<i>Papaveris Capsule</i>		557	" <i>" Composita</i>		67
<i>Papavero</i>		557	" <i>Ichthyol</i>		677
" <i>" "</i>		555	" <i>Iodi et Amyli</i>		671
" <i>" "</i>		1082	" <i>Londinensis</i>		1086
" <i>" "</i>			" <i>Lubricans</i>		35
" <i>" "</i>			" <i>Resorcinol Potium</i>		1010
" <i>" "</i>			" <i>" Mitis</i>		1010
" <i>" "</i>			" <i>Urina</i>		1248
<i>Paracetamol Hydrochloride</i>		415	" <i>de Vena</i>		941
<i>Paracotona</i>	2 to 3 gr	450	" <i>Zinci Chloridi</i>		1244
" <i>" "</i>		44	" <i>" "</i>		1244
" <i>" "</i>		42	" <i>" "</i>		1248
" <i>" "</i>			" <i>" "</i>		1010
" <i>amidine</i>		416	" <i>" "</i>		125
<i>Paraffin Injection</i>		558	" <i>" "</i>		1177
" <i>Oxygenated</i>		564	" <i>" "</i>		139
<i>Paraffina Molles</i>		562	" <i>" "</i>		495
<i>Paraffina</i>		555	" <i>" "</i>		498
<i>Paraffinum</i>		555	" <i>" "</i>		472
" <i>Carbotatum</i>		34	" <i>" "</i>		87
" <i>Datum</i>		555	" <i>" "</i>		780
" <i>Liquidum</i>		560	" <i>" "</i>		20
" <i>Massa</i>		563	" <i>" "</i>		
" <i>Molle</i>		562	" <i>" "</i>		
" <i>Solidum</i>		558	" <i>" "</i>		
" <i>" "</i>		545	" <i>" "</i>		
" <i>" "</i>		545	" <i>" "</i>		
" <i>" "</i>		565	" <i>" "</i>		
" <i>" "</i>		565	" <i>" "</i>		
<i>Paraldehyde</i>	1 to 2 fl drim	565	" <i>" "</i>		
<i>Para Monochlorophenol</i>		36	" <i>" "</i>		

Official Names in Roman, all others in Italics			PIE		
	Dose	Page		Dose	Page
<i>Pastillus Acidi Carbonici</i>		95	<i>Petrolatum Molle</i>		862
" <i>Iconia</i>		95	" <i>Spissum</i>		865
" <i>Apomorphina et Codeina</i>		180	<i>Petrolatum Benzin</i>		237
" <i>Isosmuthi</i>		246	" <i>Pther</i>	237	864
" <i>et Morphina</i>		246	" <i>Spiritus</i>		864
" <i>Cocaine</i>		412	<i>Petrosulfat</i>		696
" <i>et Morphina</i>		412	" <i>de Longona</i>		906
" <i>Codeina</i>		41	" <i>Isferminol</i>		764
" <i>Iodoforma</i>		661	" <i>Isarbitis Nil</i>		708
" <i>Ipecacuanha cum Opio</i>		77	" <i>Phellandrium</i>		499
" <i>Menthol</i>		772	" <i>Phenacitimum</i>	5 to 10 gr	87
" <i>Morphina Hydrochlorici</i>		777	" <i>cum Caffena Isferescens</i>	60 to 120 gr	87
" <i>Sulphuris Compositus</i>		1177	" <i>Isenatium</i>	5 to 10 gr	8
" <i>de Fuhon Officinale</i>		19	" <i>Isenat de Soude Solution</i>		8
" <i>de Fuhon Officinale</i>		19	" <i>Phenacitimum</i>	5 to 10 gr	877
" <i>Isenat</i>		8	" <i>Isferescens</i>	60 to 120 gr	870
" <i>Isenat Solution</i>		368	" <i>Phenacitium Hydrochloridum</i>	5 to 10 gr	876
" <i>Isenat Oil</i>		827	" <i>Salicylas</i>		877
" <i>Isenat Lactey</i>		601	" <i>Phenol</i>		27
" <i>Pearson's Antiseptic</i>		1	" <i>Isenol Camphor</i>		36
" <i>Cerate</i>		918	" <i>Isolation</i>		36
" <i>Isenat de Longona</i>		100	" <i>Isolation</i>		36
" <i>Isenat Catu</i>		18	" <i>Isolation</i>		36
" <i>Isenat</i>		973	" <i>Isolation</i>		36
" <i>Pelliturina</i>		578	" <i>Isolation</i>		36
" <i>Pelliturina</i>		578	" <i>Isolation</i>		36
" <i>de Panret</i>		579	" <i>Isolation</i>		36
" <i>Pelliturina Sulphas</i>		579	" <i>Isolation</i>		36
" <i>Tannicus</i>	6 gr	779	" <i>Isolation</i>		36
" <i>Pelliturina</i>	5 to 10 gr	972	" <i>Isolation</i>		36
" <i>Pentol</i>		75	" <i>Isolation</i>		36
" <i>Pentamulfure d Andimome</i>		171	" <i>Isolation</i>		36
" <i>Pepi Cubebe</i>		462	" <i>Isolation</i>		36
" <i>Pepi Negro</i>		902	" <i>Isolation</i>		36
" <i>Iepione</i>		323	" <i>Isolation</i>		36
" <i>Iepi, Black</i>		10	" <i>Isolation</i>		36
" <i>Pipermint Oil et</i>		764	" <i>Isolation</i>		36
" <i>Isenat Medici</i>		868	" <i>Isolation</i>		36
" <i>Isenat</i>	5 to 10 gr	868	" <i>Isolation</i>		36
" <i>Saccharatum</i>		872	" <i>Isolation</i>		36
" <i>Peptonised Beef Jelly</i>		856	" <i>Isolation</i>		36
" <i>Gruel</i>		856	" <i>Isolation</i>		36
" <i>Milk</i>		856	" <i>Isolation</i>		36
" <i>Milk Gruel</i>		856	" <i>Isolation</i>		36
" <i>Nutritive Enemata</i>		856	" <i>Isolation</i>		36
" <i>Peptonising Powder</i>		856	" <i>Isolation</i>		36
" <i>Powders and Tablets</i>		856	" <i>Isolation</i>		36
" <i>Percha Laminata</i>		589	" <i>Isolation</i>		36
" <i>Perchloride of Mercury</i>		621	" <i>Isolation</i>		36
" <i>Percolation, process of</i>		1215	" <i>Isolation</i>		36
" <i>Perhydrol</i>		640	" <i>Isolation</i>		36
" <i>Periodotetrahydroparamethyl</i>		1202	" <i>Isolation</i>		36
" <i>oxydoholium</i>		973	" <i>Isolation</i>		36
" <i>Perire de Africa</i>		97	" <i>Isolation</i>		36
" <i>de Caucasus</i>		960	" <i>Isolation</i>		36
" <i>Permanganas Kalicus</i>		960	" <i>Isolation</i>		36
" <i>Permanganate de Potassium</i>		960	" <i>Isolation</i>		36
" <i>Permanganate Potassia</i>		960	" <i>Isolation</i>		36
" <i>Peromni</i>		740	" <i>Isolation</i>		36
" <i>Peru Balsam</i>		114	" <i>Isolation</i>		36
" <i>Peruiche Officinale</i>		1255	" <i>Isolation</i>		36
" <i>Pessary Bases</i>		556	" <i>Isolation</i>		36
" <i>Pessary Acidi Tannini</i>		87	" <i>Isolation</i>		36
" <i>Alumini</i>		127	" <i>Isolation</i>		36
" <i>et Zinci</i>		127	" <i>Isolation</i>		36
" <i>Atropina</i>		204	" <i>Isolation</i>		36
" <i>Comina</i>		441	" <i>Isolation</i>		36
" <i>Isodupli</i>		498	" <i>Isolation</i>		36
" <i>Isodupli</i>		916	" <i>Isolation</i>		36
" <i>et Atropina</i>		916	" <i>Isolation</i>		36
" <i>Quinnia</i>		994	" <i>Isolation</i>		36
" <i>Zinc</i>		1215	" <i>Isolation</i>		36
" <i>Quinnia</i>		1215	" <i>Isolation</i>		36
" <i>Petit Grain, Oil of</i>		211	" <i>Isolation</i>		36
" <i>Isodupli</i>		237	" <i>Isolation</i>		36

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	Dose	Page		Dose	Page
<i>Pigmentum Acid Chromici</i>		59	<i>Pilula Casarea Composita</i>		340
" <i>Chloral Camphoratum</i>		364	" " <i>et Belladonna et Nu-</i>		
<i>Chrysarobini</i>		380	" " <i>cus Vomica</i>	1 to 3	341
<i>Iodo</i>		671	" <i>Cathartica Composita</i>		446
" <i>cum Aconiti Folio</i>		672	" " <i>Vegetabiles</i>		446
" " <i>Mite</i>		672	" <i>Colema Composita</i>		442
" " <i>Carbolicum</i>		36	" <i>Colchici et Hydrargyri</i>		431
" " <i>Carbolisati</i>		36	" " " <i>Composita</i>		
" " <i>Carbolizatum</i>		672	" " " <i>Compositum</i>		431
" <i>Oleatum</i>		672	" <i>Colocynthis Composita</i>	4 to 8 gr	435
<i>Iodoforini</i>		664	" " <i>et Hydrargyri</i>		436
" <i>Compositum</i>		663	" " " <i>Hysciani</i>	4 to 8 gr	433
<i>Mandi</i>		672	" <i>Conia Composita</i>		449
<i>Menthol</i>		772	" <i>Copasbi</i>		448
" <i>Mentholis Compositum</i>		772	" <i>Cycosoti</i>		344
<i>Phenol Iodati</i>		36	" <i>Cynoglossi</i>		472
" <i>Picus cum Iodo</i>		672	" " " <i>a</i>		473
" " <i>Lupuli</i>		909	" " " <i>a</i>		473
" <i>Resorcin</i>		1010	" " " <i>(Baillie & Pili)</i>		478
<i>Pilulas de Acibar</i>		122	" " <i>et Hydrargyri Com-</i>		
" <i>Androsan</i>		305	" " <i>posita</i>		478
" <i>Azules</i>		604	" " " <i>Opio Composita</i>		
" <i>de Bland</i>		513	" " " <i>(Henn & Pili)</i>		478
" <i>Bonico</i>		305	" <i>Flatera Composita</i>	1 or 2 pills	482
" <i>Cynoglossa</i>		475	" <i>Ergotini</i>		485
" <i>Pacgens</i>		838	" <i>Eunymum et Casarea</i>		601
" <i>de Vallet</i>		513	" <i>Ferrate Blandin</i>		513
<i>Pillule de Bland</i>		513	" <i>Ferri</i>	5 to 15 gr	512
" " <i>Carbonato Ferroso</i>		513	" " <i>Arsenicale</i>		509
" " <i>Vallet</i>		513	" " <i>Carbonatis</i>		513
<i>Pills, Coatings for</i>		887	" " <i>Carbonis</i>		513
" <i>Excerpts for</i>		887	" " <i>Blandin</i>		513
<i>Pilocarpina Hydrochloridum</i>	$\frac{1}{10}$ to $\frac{1}{2}$ gr	885	" " <i>Composita</i>		513
" <i>Nitras</i>	$\frac{1}{10}$ to $\frac{1}{2}$ gr	893	" <i>Iodidi</i>		519
" <i>Phenas</i>		896	" " <i>Quinnæ et Strych-</i>		
" <i>Salicylas</i>		896	" " <i>nini Phosphatis</i>		529
<i>Pilocarpine</i>		895	" " <i>Iodidi Com-</i>		
<i>Pilocypus</i>		685	" <i>posita</i>		1255
<i>Pilule (group)</i>		897	" <i>Ferrici Phosphatis cum</i>		
<i>Pilula Aloes Barbadosis</i>	4 to 8 gr	121	" <i>Quinnæ et Strychnini</i>		529
" " <i>et Belladonnæ</i>		123	" <i>Galbani Composita</i>	4 to 8 gr	549
" " <i>Composita</i>		123	" <i>Gutta Aloetica</i>		124
" " <i>diulula</i>	5 or 4 gr	123	" <i>Hydrargyri</i>	4 to 8 gr	604
" " <i>et Asafetide</i>	4 to 8 gr	122	" " <i>Carbolici</i>	2 to 4 pills	607
" " <i>Ferri</i>	4 to 8 gr	122	" " <i>cum Creta et Opio</i>		634
" " <i>Jalap</i>		123	" " " <i>Ipeca-</i>		
" " <i>Mastiches</i>		123	" " " <i>cuanha</i>		643
" " <i>Mithri</i>	4 to 8 gr	122	" " <i>Iodidi Vindis</i>		613
" " <i>Mithri Com</i>		123	" " <i>cum Opio</i>		604
" " <i>Podophylli Com-</i>			" " <i>Alleo</i>		605
" " <i>posita</i>		926	" " <i>Sub Chloridi Com-</i>		
" " <i>Socotriniæ</i>	4 to 8 gr	122	" " <i>posita</i>	4 to 8 gr	630
" <i>Aloeticæ</i>		123	" " <i>et Jalapa</i>		650
" " <i>Com</i>		123	" " " <i>Sam-</i>		
" " <i>posita</i>	12, 926		" " <i>moni</i>		630
" " <i>Strychnina Com-</i>		124	" <i>Ipecacuanhe et Scilla</i>	4 to 8 gr	657
" " <i>posita</i>		131	" " <i>et Utrinea</i>	4 to 8 gr	690
" <i>Asafoetida</i>		123	" <i>Jalapin</i>		705
" " <i>Composita</i>	See Pi-	198	" <i>Kali Iodati</i>		957
" " <i>lula Galbani</i>	Comp		" <i>Lazarini Composita</i>		124
" <i>Asiatica</i>		549	" <i>Mallorum</i>		1006
" <i>Atropina</i>		18	" <i>Moschi</i>		797
" " <i>et Morphine</i>		204	" <i>Myrrha Ferrata</i>		513
" <i>Atropinæ</i>		204	" <i>Opio</i>		885
" <i>Atropinæ</i>		513	" <i>Phosphori</i>	1 to 2 gr	885
" <i>Atropinæ</i>		513	" " <i>Composita</i>		885
" <i>Atropinæ</i>		513	" " <i>c Quinnæ</i>		885
" <i>Atropinæ</i>		513	" " <i>c Sero</i>		909
" <i>Atropinæ</i>		513	" <i>Picus</i>	2 or 3 pills	913
" <i>Atropinæ</i>		513	" <i>Phumhi Opio</i>		
" <i>Atropinæ</i>		513	" <i>Podophyllini, Belladonna</i>	2 to 4 gr	926
" <i>Atropinæ</i>		513	" " <i>et Capsici</i>		
" <i>Atropinæ</i>		513	" " <i>et Belladonna et Nu-</i>		
" <i>Atropinæ</i>		513	" " <i>cus Vomica</i>		926
" <i>Atropinæ</i>		513	" <i>Composita</i>		926
" <i>Atropinæ</i>		513	" <i>Quinnæ Sulphatis</i>	2 to 8 gr	1001

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	Do c	Page	Do c	Page
Lithia Omnis Sulphatis Composita		1006	Plumbi Acetis	911
" " cum Laro		1006	" Carbonis	915
" Theri et Colocynthis et Hydnangii		1016	" Iodidum	915
" " Composita	4 to 8 m	1016	" Ictas	60
" " et Vitis Lemna		1014	" Olea	918
" Saponis Composita	2 to 4 m	88	" Oxidum	916
" Scammonei Composita	4 to 8 m	1060	" Subacutatis Liquor Fortis	919
" Scilla Composita	4 to 8 m	1064	Plumbum	910
" Scirrhulana Chia		1198	" Actuum	912
" Prunum Phosphatum		51	" " Iusium Solutum	919
" " Ialavanatum		1255	" " Depuratum	912
" Urogenis Composita		1064	" Carbonium	914
" Valeriana Composita		1255	" " Basiuum	914
" Vallet		613	" Iodatum	915
" Vallet		613	" Oxidatum	917
" Vinet et Belladonna		1248	" Subacutum Solutum	919
" " Orati et Belladonna		1248	Plumbei s Pill	630
" " Valerianalis		1255	Pneumonia	71 to 30 m
" Zittmann		610	Pneumococcus Antitoxin	126
Lithia Apurantes Stahlii		123	Po de Canella Comp	598
Lithia de Aloes et Communituta		305	" Ipecacuanha Composita	648
" Carbonato Lirioso		513	" Oxidum de Citrato de Magnesia	752
" Cynoglossa		838	" " Magnesia Gasea	752
" Mercurialis		604	" Iodoformum	923
" de Opio Comp		838	" Iodatum	922
Pilule Anderson		305	" Podophylli Indica Lesina	922
Pilule de Aloes et de Saxon		122	" " Alia omnia	922
" Carbonate Liriosa		513	" " Rerum	923
" " de Fer Composita		513	" " Alia omnia	922
" " de Cynoglossa Opacis		472	" Podophyllin	923
" " Iodum Mercurialis Opacis		61	" Podophyllinum	923
" Mercurialis Simplex		604	" Podophyllotomum	924
" " de Podophylline Ictas		921	Poison Poi	1018
" " de Liriod		613	Povere Nov	902
Pimenta	10 to 30 m	898	" " de Guine	323
" " da Jamaica		898	" " de Bourgogne	906
Pimentao		824	" " Root	892
Pimenta de la Jamaica		806	" " Root	892
" " Gord		898	" " Root	892
" " Negra		902	" " Root	892
Pimento de Indias		823	" " Root	892
Pimprilla		165	" " Root	892
Pini Oleum	1 to 5 min	900	" " Root	892
" " Sylvestris Oleum		901	" " Root	892
Pinol		900	" " Root	892
Prambi		910	" " Root	892
Piperi Nigrum		902	" " Root	892
Piperazine	5 to 15 m	904	" " Root	892
" " Quinque	5 to 15 m	905	" " Root	892
Piperidine		903	" " Root	892
" " Acid Tartrati	10 to 15 m	904	" " Root	892
" " Guaiaculati	5 to 30 m	904	" " Root	892
Piperina	3 m	903	" " Root	892
Picro		972	" " Root	892
Picrogallolo		72	" " Root	892
Picrostema		975	" " Root	892
Piscidia		906	" " Root	892
Pissentit		1193	" " Root	892
Pitch, Black		910	" " Root	892
" " Burgundy		906	" " Root	892
" " Plaster		906	" " Root	892
Pix Burgundica		906	" " Root	892
" " Carbonis Preparata		907	" " Root	892
" " Liquida	5 to 10 min	908	" " Root	892
" " Lithanthrasia		907	" " Root	892
" " Solida		908	" " Root	892
Plaue Vaccina		1268	" " Root	892
Plaster Muls (Unna)		589	" " Root	892
" " of Paris		294	" " Root	892
Plasters See Emplastia			" " Root	892
Platre Coalit		907	" " Root	892
Plomb		910	" " Root	892
Plomo		910	" " Root	892

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	Dose	Pt.		Dose	Pt.
<i>Pomerianzenschale</i>		206	<i>Præcipitatum Album</i>		651
<i>Pommade Acide Borique</i>		20	<i>Præcipitatum Chalk</i>		280
" <i>Belladonna</i>		25	" <i>Sulphur</i>		1176
" <i>de Calomel</i>		650	<i>Præcipitatum Album</i>		628
" <i>Camphire</i>		512	<i>Præcipit. blanc</i>		496
" <i>de Chloroforme</i>		77	<i>Prepared Chalk</i>		625
" <i>Lipostaphure Intum</i>		521	<i>Preservative Solution for Ana-</i>		
" <i>" Lerte</i>		521	<i>tomical Subjects</i>		
" <i>de Goudron</i>		909	<i>Proof Spirit</i>		1199
" <i>d'Indure de Potassium to-</i>			<i>Propylamine</i>		1111
<i>lu</i>		670	" <i>Rhynchostomide</i>		1219
" <i>Mercurielle Laible</i>		604	<i>Protargol</i>		188
" <i>Naphthole</i>		604	<i>Proto Indure de Mercure</i>		611
" <i>d'Oxyde de Mercure Intum</i>		619	<i>Proto Induratum Hippocratice</i>		61
" <i>de Phenol</i>		4	<i>Protodurum et de Manganese</i>		788
" <i>Sanguifate de Phenyl</i>		1059	" <i>" Mercure pur</i>		617
" <i>" Styren</i>	1005, 1171		" <i>Præcipitatum</i>		
" <i>Sulphureuse</i>		933	" <i>de Mercure Induratum</i>		617
<i>Pommades</i>		1221	<i>Protosulfo di Pombo</i>		916
<i>Pondus Extract</i>		106	<i>Purum Vermine Cortex</i>		969
<i>Pondus Pill</i>		926	<i>Purum</i>		910
<i>Popp, Chin-ic</i>		857	<i>Purissim. of Potash, Yellow</i>		930
<i>Purissim. Chin</i>		709	<i>Pseudocutane</i>		96
<i>Potassa Caustica</i>		927	<i>Pseudovine</i>		1231
" <i>" 205 la Cal</i>		928	<i>Pseudopelletierine</i>		76
" <i>" Prusa</i>		928	<i>Pseudopurine Granatatum</i>		576
" <i>cum Calce</i>		928	<i>Psychotria</i>		602
" <i>Sulphurata</i>	1 to 5 gr	931	<i>Psych. Semen</i>		971
<i>Potassi Acetas</i>	10 to 60 gr	933	<i>Ptyocarpus Lignum</i>		971
" <i>Benzas</i>	15 to 20 gr	935	<i>Pulpe de Cassie</i>		345
" <i>Burcharis</i>	5 to 30 gr	935	<i>Pulcatilla</i>		973
" <i>Bichonias</i>	15 to 2 gr	937	<i>Pulv. Phenolom Comp</i>		879
" <i>Bitartras</i>		966	<i>Pulvis Acidi Borici Compositus</i>		26
" <i>Bromidum</i>	5 to 30 gr	939	"		77
" <i>Carbonas</i>	5 to 20 gr	942	" <i>Aloes et Canellu</i>		1090
" <i>Cantharidin</i>		822	"		124
" <i>Chil-ic</i>	5 to 15 gr	944	"		15
" <i>Citras</i>	10 to 40 gr	947	"		573
" <i>Effet resens</i>		948	"		170
" <i>Cyanidum</i>		949	" <i>Antimonidis</i>		
" <i>Ferrocyanidum</i>		950	" <i>Atomiticus, vel Pulv. Chin</i>		391
" <i>Guaiacol Sulphonas</i>		557	" <i>nam Comp</i>		291
" <i>Hydras</i>		927	" <i>Compositus</i>		610
" <i>Hydroxidum</i>		928	" <i>Basilicus</i>		61
" <i>Hypophosphis</i>		951	" <i>Calomelanos et Acidi Borici</i>		61
" <i>Iodidum</i>	5 to 20 gr	951	" <i>" Annul</i>		61
" <i>Nitras</i>	5 to 20 gr	957	" <i>" Amis Oculi</i>		61
" <i>Osmas</i>		67	" <i>cum Rhco</i>		61
" <i>Orychinolus Sulphonas</i>		901	" <i>Calchu Compositus</i>	10 to 40 gr	347
" <i>Purissim. Chin</i>	1 to 5 gr	900	" <i>Chinuinom Compositus</i>	10 to 40 gr	395
" <i>et Sicut Lact</i>		1088	" <i>Citric Atomiticus</i>	10 to 60 gr	457
" <i>Soromodolas</i>		1138	" <i>" c Opio</i>	10 to 40 gr	457
" <i>Sulphur</i>	10 to 40 gr	963	" <i>Compositus</i>		457
" <i>Sulphocarbolas</i>		97	" <i>Cynoglossa Compositus</i>		472
" <i>Lutras</i>	50 to 240 gr	964	" <i>Boycu</i>		685
" <i>Acidus</i>	20 to 60 gr	966	" <i>Effer resens Laxans</i>		1090
<i>Potissio Taitate of Antimony</i>		173	" <i>Elatini Compositus</i>	1 to 4 gr	48
<i>Potassium</i>		927	" <i>Glycyrrhiz Compositus</i>	60 to 100 gr	173
" <i>Cantharidate</i>		522	" <i>" c</i>	20 to 40 gr	585
" <i>Dichromate</i>		957	" <i>" c</i>		1016
" <i>Guaiacol Sulphonate</i>		557	" <i>" Subchlorus Com-</i>		
" <i>Hydrate</i>		927	<i>positus</i>		610
" <i>Hydrogen Carbonate</i>		935	" <i>T</i>		614
<i>Potio L</i>		948	" <i>"</i>	5 to 15 gr	685
" <i>" c</i>		702	" <i>sue Fmetuna</i>		67
" <i>resens</i>		702	" <i>Opatus</i>		685
" <i>Magnesi Citric Aero-</i>		752	" <i>et Opn</i>		685
" <i>phora</i>		948	" <i>Thebaicus</i>		685
" <i>Rupen</i>		947	" <i>Valip Compositus</i>	20 to 60 gr	703
<i>Potion Gazeuse</i>		947	" <i>Kaladane Compositus</i>	20 to 60 gr	703
" <i>Gommeuse</i>		4	" <i>Kino</i>	5 to 20 gr	711
<i>Poudre contre la Coque</i>		772	" <i>pro Lacte Humano</i>		854
" <i>d'Ipeacuanha Opio-re</i>		688	" <i>Liquorista Compositus</i>		573
" <i>de Sulfur Composit et</i>		625	" <i>Lobelia</i>		741
" <i>u. de Turp. juce</i>		625	" <i>Magnesi Borocitratis</i>		20
			<i>Compositus</i>	30 to 60 gr	20

QUI		Official Names in Roman, all others in Italics	
	Dose	Page	
Quinine Bihydrochloride		995	<i>Resorcin Monoacetat</i>
" Formate, Basic		984	<i>Resorcinol</i>
" " Neutral		984	" <i>Phthalen Anhydride</i>
Quinoidin	5 to 15 gr	990	<i>Resorcinarum</i>
Quinolone		991	<i>Respirators</i>
" Bismuth Sulphocyanide		991	<i>Rhamni Frangula Cortex</i>
Quinosol		991	<i>Rhabarber</i>
Quinolopine		517	<i>Rhamni Purshiani Cortex</i>
Quinquina Jaune		85	<i>Rhizomy Root</i>
" Rouge	82	1257	" and <i>Cocaine Lozenges</i>
			" <i>Lozenges</i>
<i>Rabano Rusticano</i>		191	<i>Rhei Revlix</i>
<i>Rabao Rustica</i>		191	{ 1 to 10 gr (repeated) 15 to 30 gr (single) }
<i>Rabobarbo</i>		1012	
<i>Racine d'Actee Groupes</i>		80	<i>Rheumatine</i>
" <i>d'Anigue</i>		192	<i>Rhi onia (Giamius)</i>
" <i>de Belladone</i>		228	<i>Rheo</i>
" " <i>Colombo</i>		295	<i>Rhizos Petali</i>
" " <i>Gentiana</i>		561	<i>Rhododol</i>
<i>Racine de Belladonna</i>		928	<i>Rhizobul Root</i>
<i>Radium</i>		1222	<i>Rhizobul de Chine</i>
" <i>Hydro</i>		1222	<i>Rhus Aromaticum</i>
" <i>Sulphate</i>		1223	" <i>(Habra)</i>
<i>Rafano Rusticano</i>		191	" <i>Toxicodendron</i>
<i>Rafort</i>		191	<i>Rien</i>
<i>Ranz de Belladonna</i>		228	<i>Ricini Oleum</i>
" " <i>Colombo</i>		295	<i>Ricinusol</i>
" " <i>Peltie</i>		972	<i>River seche Tank</i>
<i>Ratanha</i>		713	<i>Rizoma de Arnica</i>
<i>Ratanhia</i>		713	<i>Rochelle Salt</i>
" <i>du Perou</i>		713	<i>Rohrenkassie</i>
<i>Ratanhiawurzel</i>		713	<i>Romera</i>
<i>Ratma</i>		713	<i>Romische Berthamunzel</i>
<i>Rautenol</i>		1029	" <i>Kamille</i>
<i>Reagents used in chemical testing</i>	1200 to 1291		<i>Romschmanzol</i>
<i>Rectified Spirit</i>		1146	<i>Rosmarini Essentia</i>
<i>Red Bone Marrow</i>		761	<i>Rosa Roja</i>
" <i>Gum</i>		492	<i>Ros e Aqua</i>
" <i>P</i>		1017	" <i>Gallia Petali</i>
" <i>P</i>		620	" <i>Oleum</i>
" <i>Rose Petals</i>		1021	<i>Rose Rouge</i>
" <i>Sandal Wood</i>		971	<i>Rosenol</i>
" <i>Sanders Wood</i>		971	<i>Rosencasser</i>
<i>Reduced Iron</i>		534	<i>Rosmarini Oleum</i>
<i>Reductores Fusen</i>		534	<i>Rosolaccio</i>
<i>Regale</i>		570	<i>Rosolic Acid Solution</i>
<i>Reglose</i>		570	<i>Rubidium Ammonium Linnide</i>
<i>Regulate Amalgam Mixture</i>		578	" <i>Linide</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Jodide</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Rubini's Essence of Camphor</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Rubarbo</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Rumex</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Ruta Oleum</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sabao, Animal</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Vegetal</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Sabina Catemont</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sabine</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sabina</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sabinqueno</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Saccharum</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharated from Carbonate</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Solution of Lime</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharin Dose</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharinum</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharinum</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Soluble</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharinum Carapen</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Chetia</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Lutis</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	" <i>Purificatum</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacharine de Catharine (P)</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>near</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sacred Luk</i>
<i>Regulate Silver Waxes and Spirits</i>		1150	<i>Sedebamus et al</i>

Official Names in Roman, all others in Italics			SCI	
	Dose	Page		Dose
Saffron <i>See Crocus</i>		457	<i>Sanguisuela</i>	597
Safran		457	<i>Sanguisuga</i>	597
Safron		10-8	<i>Sanguis Bovinus Haemecatus</i>	593
Sagradarinde		337	<i>Santas Disinfectants</i>	1198
Sajodin		673	<i>Sanoform</i>	554
Salacetol		78	Santalal Oleum	5 to 30 min 1041
Sal Alembroth		626	<i>Santalum Indicum</i>	971
<i>Diomedium F. fl. nescens</i>		912	Santonium	2 to 5 gr 1044
<i>Carolinum F. fl. nescens</i>		1132	<i>Santyl</i>	1044
<i>Prunum F. fl. nescens</i>		1090	<i>Sapo</i>	1049
<i>Polychestum</i>		965	<i>Albissimus Diagonistatum</i>	1047
<i>Polychestum</i>		963	<i>Albus Oleaceus</i>	1049
<i>Segnette</i>		10-8	<i>Amurhis</i>	1047
<i>Prunella</i>		959	<i>Butyraceus</i>	1047
<i>Volatide, Spirit of</i>		141	<i>Durum</i>	5 to 15 gr 1048
Salt p		1035	<i>Hispanicus Albus</i>	1049
Salt Misi		1035	<i>Salpinx</i>	705
Salicidato de litina eferescite		739	<i>Kalmus</i>	1051
<i>de Fenol</i>		1037	<i>Venalis</i>	1051, 1054
Salicylate de Naphthyle B		806	<i>Medicatus</i>	1047
<i>Neutre de Sodium</i>		1126	<i>Medicinalis</i>	1047
<i>de Phenyle</i>		1037	<i>Mollis</i>	1051
Salicidato de Ictio		739	<i>Officinalis</i>	1049
<i>Sodium</i>		1126	<i>Oleaceus</i>	1049
Salicinium	5 to 20 -1	1035	<i>Sebacinus</i>	1047
Salicylas Natruo Coffeius		277	<i>Stearinus</i>	1047
<i>Natruo cum Coffeino</i>		277	<i>Superadipatus</i>	1051
Salicylated Vasoliment		78	<i>Venetus</i>	1049
Salicylic Acid		73	<i>Verris</i>	1051
<i>Collodion, Compound</i>		77	<i>Sapone Animale</i>	1047
<i>and Creosote I luster Mulls</i>		77	<i>Medicinalis</i>	1048
<i>Collodion</i>		77	<i>de Potassa</i>	1051
<i>Cotton</i>		77	Saponification value of fixed oils	1317
<i>Dressing</i>		77	Saponification value of Gums	
<i>Gauze</i>		77	<i>and Gum resins</i>	1318
<i>Jute</i>		77	Saponin	980
<i>Jute</i>		77	Saponis Emplastrum Camphor	
<i>Ointment</i>		76	<i>atum</i>	1050
<i>Wool</i>		77	Sapotoxin	980
Salicylguanine Salicylate		985	Sappan	1054
Salicylsare		73	Saprol	44
Salicyllol		73	Sarsapareille de Mexique	1054
Saligenin	4 gr	1037	Sarsaparilla	1054
Salinigrin		1037	Sassa Radix	1054
Salpinx	10 to 30 gr	880	Sassafras, Black	1059
Salt		305	<i>Medulla</i>	1050
Salatanal		79	<i>Oil of</i>	1059
Salt Nigra		1037	<i>Radix</i>	1058
Salicoll	15 to 30 gr	577	Sassy Bark	492
Salicreol	6 to 20 -1	455	Sauco	1040
Salol	5 to 15 -r	1037	Saures Channidhydriochtonid	994
<i>Camphor</i>		1039	Savin Tops	1030
<i>Mouth Wash</i>		1039	Savin Animal	1047
<i>Varnish</i>		1040	<i>Medicinal</i>	1048
Salophen	10 to 30 -r	1040	<i>Nom</i>	1051
Salopguanine	15 to 30 -1	987	Scammomul I ulis	1050
Saltpetresare		61	<i>Resin</i>	5 to 8 gr 1060
Salt of Ictio		942	Scammomul	705
<i>Wisdom</i>		625	Scammomul	5 to 10 gr 1061
Saltpetre		957	Schierling	437
Salts Anti catarrhal		4	Schlangenwurzel	380
Salutibath		56	Schulze's Solution	1344
Salumun		129	Schuster's Prestille	97
Salv. Mulls (Umm)		100	Schnefellebe	932
Salvo Petrolia		862	Schnefellebe	70
Salvamine		50	Schnefellebe	83
Sambucus Flores		1040	Schnefellebe	96
Sambuco		1040	Schwere gebannte Magnesia	747
Sambucum		1040	Schill	1 to 2 gr 1062
Sandil wood, Oil of		1041	Schillam	1062
<i>Oil</i>		971	Schille	1062
Sandalo Rind		971	Schillipin	1062
Sandarach Varnish		898	Schillipin	1062
Sandelol		1041	Schillipin	1062
Sandels Wood 1 ed		971	Schillipin	1062
Sandolo Rind		971	Schillipin	1062
Sandolo		597	Schillipin	1062

SCI	Official Names in Roman, all others in Italics				
	Dose	Page		Dose Page	
		1075	<i>Sirup d Ethio</i>		107
		489	" <i>de Baume de Toli</i>		220
Scoparin Cacumini		1066	" " <i>Gomme</i>		4
Scopola		1068	" " <i>Goudron</i>		910
Scopolamina		648	" <i>Iodotannique</i>		672
" "		653	" " <i>Phosphate</i>		672
" "		649	" <i>de Mures</i>		776
" <i>Hydrochloridum</i>		653	" " <i>Quinquina</i>		392
<i>Scorza di Lemone</i>		722	" " <i>Infusit Compose</i>		192
<i>Scotch Paragoric</i>		811	" <i>Ichthyol Compose</i>		101
<i>Scott's Dressing</i>		601	" " <i>Sarsaparilla Compo</i>		1055
<i>Scutellaria</i>		1060	<i>Sirupus Feni Potatis Compo</i>		
<i>Scutellarin</i>	1 to 5 gr	1069	" "		506
<i>Sebo</i>		1078	" <i>Puris cum Codomo</i>		910
" <i>de Carneio</i>		1075	" <i>Senna cum Manni</i>		107
<i>Sebum Saponatum</i>		77	<i>Slaked Lime</i>		286
<i>Secale cornutum</i>		485	<i>Slippery Elm</i>		1221
<i>Sejala Coimuta</i>		48	<i>Snake root</i>		1077
<i>Seulebasteride</i>		775	<i>Sodium</i>		1106
<i>Scutitz Powder</i>		1090	<i>Soap Bark</i>		980
<i>Scifemide</i>		940	" <i>Curd</i>		1017
<i>Sciles & Antisplac</i>		261	" <i>Hard</i>		1018
<i>Sci de la Sagesse ou de la senn</i>		625	" <i>Plaster</i>		1050
<i>Semen Calabariense</i>		886	" <i>Salt</i>		1011
" <i>Myrsine</i>		795	<i>Soapstone</i>	120	119
" <i>Physostigmatis</i>		886	<i>Socotrine Aloes</i>		118
" <i>Strychni</i>		811	<i>Socotrinische Aloe</i>		118
<i>Semence de Colchique</i>		429	<i>Soil & Alum</i>		125
<i>Semi de Colchico</i>		429	" <i>Cauteria</i>		1086
<i>Semilla de Colquico</i>		429	" <i>Tinctura</i>	120 to 240 gr	1088
<i>Semina Calabar</i>		886	<i>Sodin Acetas</i>		1090
<i>Sen de Espana</i>		1072	" <i>Amias</i>	5 to 15 gr	160
<i>Senape Nera</i>		1079	" <i>Arenas</i>	1 to 1 gr	1092
<i>Sene</i>		1074	" <i>Lenzoas</i>	5 to 30 gr	1094
<i>Senegale Radix</i>		1070	" <i>Ribosas</i>		209
<i>Senssane</i>		1079	" <i>Rictrbonas</i>	5 to 30 gr	1096
<i>Senna</i>	10 to 20 gr	1072	" <i>Rionidium</i>	5 to 30 gr	1100
" <i>Alexandrian</i>		1072	" <i>Cacodylus</i>		1101
" <i>East Indian</i>		1073	" <i>Calbonas</i>	5 to 30 gr	1107
" <i>Tinnivelly</i>		1073	" <i>Exsiccatus</i>	5 to 10 gr	1108
<i>Senneseblatter</i>		1072	" <i>Monohydatis</i>		1107
<i>Serpentaria Rhizom</i>	10 to 15 gr	1077	" <i>Chloras</i>		947
<i>Serum for Hay Fever</i>		1204	" <i>Chloridum</i>	10 to 60 gr	1109
<i>Serum Benzoinum</i>		100	" <i>Cinnamas</i>	2 to 5 gr	1112
" <i>Benzoinatum</i>		241	" <i>Citrus</i>		1114
" <i>Phosphoratum</i>		985	" <i>Citro tart is Effavescent</i>	60 to 120 gr	1099
" <i>Preparatum</i>		1078	" <i>Dithio salicylas</i>	1 to 5 gr	1119
" <i>Salicylitum</i>		77	" <i>Ethylitis Liquor</i>		1115
<i>Sherly</i>		1234	" <i>Fluoridum</i>		46
<i>Sicco</i>		593	" <i>Glycidum</i>		40
<i>Sidonal</i>		905	" <i>Glycocholas</i>	2 to 15 gr	1116
" <i>New</i>	5 to 15 gr	905	" <i>Hydroxidum</i>		1096
<i>Silbernitrat</i>		194	" <i>Hypophosphus</i>	5 to 10 gr	1116
<i>Silberoxyd</i>		190	" <i>Hyposulphus</i>		1114
<i>Silver and Preparations See</i>			" <i>Ichthyosulphona</i>		637
" <i>Argentum</i>		184	" <i>Iodidum</i>	5 to 20 gr	1118
" <i>Albuminate</i>		188	" <i>Lactas</i>		60
" <i>Citrate</i>		188	" <i>Nitris</i>	1 to 2 gr	1117
" <i>Fluoride</i>		189	" <i>Nucleinas</i>		810
" <i>Glutin</i>		187	" <i>Oleas</i>		1051
" <i>Ichthyolate</i>		188	" <i>Panacresolinas</i>		44
" <i>Iodide</i>		187	" <i>Permannanias</i>		96
" <i>Lactate</i>		187	" <i>Phenol-sulphonas</i>		1114
" <i>Nitrate</i>		184	" <i>Picoididum</i>		815
" <i>Nucleinate</i>		810			
" <i>Oxide</i>		190	" <i>Phosphus</i>	{ 50 to 120 gr (in practice) 1 to 2 oz (single) }	1123
" <i>Protein</i>		188	" " <i>Acidus</i>	{ 50 to 60 gr (in practice) 1 to 2 oz (single) }	1126
" <i>Thiohydriocarbure sulpho</i>		188			
" <i>nate</i>		188			
<i>Simaruba</i>	15 to 30 gr	1079			
<i>Sinapis</i>		1079			
" <i>Albe Sinini</i>		1080			
" <i>Nigr Sinini</i>		1081			
<i>Sinapisem en Fluidi</i>		1082			
<i>Sinupis de Papet</i>		1082			
<i>Sirolin</i>		587			
<i>Sirup d Acide Citrique</i>		106			

SOL		Official Names in Roman, all others in Italics	
	Dose	Page	
<i>Solutio de Elixirino com (alibi)</i>			
" <i>Gaz Sulphureo</i>		188	<i>Spiritus Amosi</i> 5 to 20 min 106
" <i>Todo Todelado</i>		80	" <i>Ammoniac Compositus</i> 1 to 2 fl. drim 192
" <i>de Soda Chlorada</i>		670	" <i>Ammoniac Compositus</i>
" <i>Subacetato de Chumbo</i>		301	" <i>Cajuputi</i> 5 to 20 min 277
<i>Solutio</i>		919	" <i>Campfoi</i> 5 to 20 min 309
" <i>Alcoholis Etica di</i>		44	" <i>Compositus</i> 317
" <i>Cloruro Plumbi</i>		524	" <i>Emilia</i> 1010
" <i>Idronologica de Bulmaro</i>			" <i>Capulatus</i> 1010
" <i>di Mercurio</i>		624	" <i>Capulorum</i> 1010
<i>Solutio</i>		14	" <i>Chlorotomum</i> { 5 to 20 min (repeated) 5 to 40 min (single) } 576
<i>Somatose</i>		1138	" <i>Cinnamoni</i> 5 to 20 min 599
" <i>Iron</i>		1138	" <i>Citri</i> 726
" <i>Liquid</i>		1138	" <i>Cresoti</i> 154
" <i>Milk</i>		1138	" <i>Dulcisi</i> 1119
<i>Somnula</i>		1132	" <i>Pimentis</i> 1112
<i>Somnat</i>		1236	" <i>Gaultheria</i> 30 min 534
<i>Sottotrato di Bismuto</i>		250	" <i>Glyceritis Adriatis</i> 809
<i>Soude Cautique Liquid</i>		1086	" <i>Pumpin</i> 5 to 60 min 708
<i>Soufre Precipite</i>		1176	" " <i>Compositus</i> 708
" <i>Sublimi</i>		1178	" <i>Lividum</i> 5 to 20 min 721
" " <i>Lave</i>		1178	" <i>Mentha</i> 767
<i>Souscarbonate de Bismuth</i>		245	" " <i>Piperita</i> 5 to 20 min 767
<i>Sousnitrate de Bismuth</i>		250	" " <i>Viridis</i> 769
<i>Sozodol</i>		1138	" <i>Methylatus</i> 1145
" <i>Cotton</i>		1138	" <i>Mycia</i> 900
" <i>Gauze</i>		1138	" <i>Myristice</i> 5 to 20 min 800
<i>Spanische Rhegen</i>		317	" <i>Arcus fragrantis</i> 1 to 4 fl. drim 706
" <i>Pfeffer</i>		323	" <i>Phosphori</i> 885
<i>Spasmodic de Cantharidate de</i>			" <i>Pimenta</i> 900
" <i>Potassium</i>		322	" <i>Racilatus</i> 1146
<i>Spasmodic Vesicans</i>		322	" <i>Rosmarini</i> 1019
<i>Spasmodic Periodicum</i>	1 to 4 gr	1067	" <i>Rosmarini</i> 5 to 20 min 1029
" <i>Sulphur</i>	1 to 2 gr	1067	" " <i>Compositus</i> 1029
<i>Spasmodic Britan</i>		1319	" " <i>Compositus</i> 107
" <i>Foreign</i>		1320	" <i>Saponato-Campfoatum</i> 1034
<i>Spearmint, Oil of</i>		768	" <i>Saponatus</i> 1034
<i>Spermacet</i>		365	" <i>Saponis</i> 1034
<i>Special Tests</i>		1319	" <i>Sedum</i> 1034
<i>Spermum</i>		1139	" <i>Sinapi</i> 1034
" " <i>Compositus</i>		401	" <i>Tenuis</i> See Proof Spirit 1139
" " <i>Compositus</i>		409	" <i>Vini Cognac</i> 1132
<i>Spirit of Chloric Ether</i>		376	" " <i>Gallici</i> 1130
" " <i>Hartsorn</i>		144	" " <i>Vino</i> 1139
" " <i>Sol Volatile</i>		141	" " <i>Vino</i> 1062
<i>Spirits of the Pharmacopoeias,</i>			" " <i>Squill</i> 530
" " <i>Alcoholic Strengths</i>		1149	" " <i>Squill</i> 121
<i>Spiritus</i>		1139	" " <i>Squill</i> 121
" " <i>Phthereus Feratus</i>		524	" " <i>Squill</i> 121
" " <i>Phthereus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	106	" " <i>Squill</i> 121
" " <i>Compositus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	106	" " <i>Squill</i> 121
" " <i>Mutatus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	107	" " <i>Squill</i> 121
" " <i>Nitrosi</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	1139	" " <i>Squill</i> 121
" " <i>Ammoniac</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	133	" " <i>Squill</i> 121
" " <i>Ammoniac</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	141	" " <i>Squill</i> 121
" " <i>Compositus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	141	" " <i>Squill</i> 121
" " <i>Letidis</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	134	" " <i>Squill</i> 121
" " <i>Feneculatus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	135	" " <i>Squill</i> 121
" " <i>Ammoniac Amatus</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	135	" " <i>Squill</i> 121
" " <i>Amigdalis Amure</i>	{ 20 to 40 min (repeated) 60 to 90 min (single) }	150	" " <i>Squill</i> 121
			" " <i>Squill</i> 121

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	Dose	Page		Dose	Page
<i>Stone Root</i>		433	<i>Sulfate Neutre de Potassium</i>		986
<i>Storace liquido</i>		1170	" "		983
<i>Storax, Prepared</i>		1170	" <i>de Protoxide de Fer Officinal</i>		530
<i>Storaxne</i>	½ to 1 m	418	" "		1160
<i>Stramonie</i>		1151	" <i>Inc Officinal</i>		1249
<i>Stramonii Folia</i>	o 2 grammes	1154	<i>Sulfato Aluminium Potassio</i>		120
" <i>Scimina</i>		1157	" <i>Cuprico</i>		460
<i>Strofantio</i>		1161	" <i>de Cobre</i>		465
<i>Strontium Bromidum</i>	5 to 30 gr	1157	" <i>Eserina</i>		889
" <i>Ersucatum</i>		1158	" <i>Ferroso</i>		580
" <i>Cinnamomus</i>		1159	" <i>Magnifico</i>		752
" <i>Iodidum</i>	7½ to 15 gr	1159	" <i>Mercurico</i>		609
" <i>Lactas</i>	20 to 30 m	1159	" <i>Potassio</i>		963
" <i>Sakoylas</i>	5 to 15 gr	1160	" <i>de Potasio</i>		963
<i>Strontium Caffeine Sulphonate</i>		278	" <i>Quinico Basico</i>		996
<i>Strophanthi Scimin</i>		1161	" <i>Neutro</i>		955
<i>Strophanthum</i>	10 to 30 m	1164	" <i>de Quinina</i>		999
<i>Strophanthusamin</i>		1161	" <i>Sodico</i>		1130
<i>Stychnin</i>	1 to 10 m	1160	" <i>de Sodio</i>		1130
<i>Stychnin e Aetas</i>		1169	" <i>Zincuo</i>		1249
" <i>Hydrobromidum</i>		1169	<i>Sulphum Stibium</i>		173
" <i>Hydrochloridum</i>	10 to 20 m	1167	<i>Sulfonate</i>		1172
" <i>Meta vanadus</i>		1169	<i>Sulfure de Potasse</i>		902
" <i>Nitras</i>		1169	<i>Sulfuretum Stibium</i>		173
" <i>Sulphas</i>		1169	<i>Sulfuro Aluminium</i>		172
" <i>Valerianas</i>		1169	" <i>(tri) Potassio</i>		932
<i>Stupa Carbosata</i>		55	" <i>de Potasio</i>		932
" <i>Composita</i>		35	<i>Sulphaminol</i>		87
<i>Styptic Collod</i>		976	<i>Sulphas Chinicus</i>		999
<i>Stypticum</i>		845	" <i>Chinini</i>		999
<i>Styptol</i>		845	" <i>Kalcus</i>		963
<i>Styracol</i>		586	" <i>Natius</i>		1130
<i>Styrax Liquide Purifd</i>		1170	" <i>Quinina</i>		999
<i>Styrax Preparatus</i>		1170	<i>Sulphate of Iron</i>		294
<i>Subacetat Plumbi Liquidus</i>		919	" <i>Mercury</i>		609
<i>Subchloride of Mercury</i>		628	<i>Sulphate de Soda</i>		1130
<i>Subcutin</i>		22	<i>Sulphocarbolu ied</i>		37
<i>Sublamin</i>		608	<i>Sulphonal</i>	10 to 30 gr	1172
<i>Sublimate Gauze</i>		625	<i>Sulphonethylmethanum</i>		1174
" <i>Wood Wool</i>		625	<i>Sulphonmethanum</i>		1172
" <i>Wool</i>		625	<i>Sulphophenylas Zincicus</i>		1253
<i>Sublimatus Corrosivus</i>		621	<i>Sulphur</i>		1170
<i>Sublimed Sulphur</i>		1178	" <i>Depuratum</i>		1178
<i>Suc de Citron</i>		727	" <i>Disside, Liquefied</i>		84
<i>Suc de Mue</i>		776	" <i>Har Loton</i>		1177
<i>Succi (group)</i>		1171	" <i>Kestores</i>		1177
<i>Succinum</i>		1172	" <i>Lotum</i>		1178
<i>Succo de Limone</i>		727	" <i>Præcipitatum</i>	20 to 60 gr	1176
<i>Succus Acalypha</i>	1 to 4 fl dram	4	" <i>Sublimatum</i>	20 to 60 gr	1178
" <i>Adhatode</i>	1 to 4 fl dram	102	" <i>Crudum</i>		1178
" <i>Belladonnæ</i>	5 to 15 min	225	" <i>Lotum</i>		1178
" <i>Citri Atrichialis</i>		728	" <i>Vendale</i>		1178
" <i>Fachitus</i>		728	" <i>Washed</i>		1179
" <i>Conii</i>	1 to 2 fl dram	435	<i>Sulphurated Antimony</i>		171
" <i>Digitalis</i>	5 to 10 min	478	" <i>Lime</i>		302
" <i>Hyoscyami</i>	½ to 1 fl dram	647	" <i>Potash</i>		983
" <i>Limonis</i>		727	<i>Sulphuretum Potassii Official</i>		932
" <i>Liquiritis</i>		572	<i>Sulphuric Acid</i>		70
" <i>Mori</i>		776	" <i>Aromatic</i>		81
" <i>Scoparii</i>	1 to 2 fl dram	1066	" <i>Crude</i>		80
" <i>Taraxaci</i>	1 to 2 fl dram	1191	" <i>Diluted</i>		82
<i>Suc de Blano Officinal</i>		1038	" <i>Nordhauson</i>		79
<i>Sucrot</i>		566	" <i>Fther</i>		103
<i>Sucrose</i>		1033	<i>Sulphuris Alcalini Unguentum</i>		1179
<i>Suet, Prepared</i>		1078	" <i>Chloridum</i>		1180
" <i>Salticæ Acid</i>		77	" <i>Iodidum</i>		1180
<i>Sugar, Milk</i>		1031	" <i>Tabellæ</i>		1177
" <i>Refined</i>		1033	<i>Sulphurous Acid</i>		83
<i>Sulf de Mouton Purifd</i>		1078	" <i>Anhydride</i>		83
<i>Sulfata de Quinina neutro</i>		988	<i>Sumbul Radix</i>		1182
<i>Sulfate d Atropine</i>		201	<i>Suppositoria</i>		1133
" <i>Basique de Quinine</i>		996	" <i>Acidi Carbolicæ</i>		33
" <i>de Calcium</i>		294	" <i>Tannici</i>		86
" <i>Cuivre</i>		466	" <i>et Belladonna</i>		87
" <i>Perenne</i>		889	" <i>Morphina</i>		87
" <i>Magnesium</i>		752	" <i>c Opio</i>		87
" <i>Mercurique Basique</i>		609			

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TIN

	Dose	Page		Dose	Page
<i>Tablets of Balsam of Tolu</i>		220	<i>Tentius de Cachou</i>		348
" " <i>Cotarnina Hydrochloridum</i>		845	" " <i>Camphré Com</i>		310
<i>Tablettes de Chlorate de Potas</i>		947	" " <i>farble</i>		310
" " <i>Chlorhydrate de Cocaine</i>		412	" " <i>Cold</i>		712
" " <i>Soufre</i>		1177	" " <i>à Essence de Citron Com</i>		727
<i>Tachol</i>		189	" " <i>posce</i>		767
<i>Taka Drastase</i>	5 to 10 gr	270	" " <i>de Fève de Saint Ignace</i>		659
<i>Tala</i>		1192	" " <i>Composce</i>		980
" <i>Borated</i>		1192	" " <i>Panama</i>		390
" <i>Purified</i>		1192	" " <i>Quinquina Rouge</i>		575
<i>Tamarind Whey</i>		1193	<i>Tela Depurata</i>	5 to 15 min	1195
<i>Tamrindus</i>	4 oz	1193	<i>Terebenthinum</i>	5 to 10 gr	1197
<i>Tanato de Mercurio</i>		610	<i>Terebinthum Canadensis</i>		1198
<i>Tannalbin</i>	15 gr	57	" <i>China</i>		441
<i>Tannic Acid</i>		84	" <i>Copahiba</i>		717
" <i>Wool</i>		87	" <i>Laricus</i>		717
<i>Tannigen</i>	1 to 10 gr	88	" <i>Veneta</i>		717
<i>Lannin</i>		84	<i>Terebinthum Oleum</i>	2 to 10 min	1198
" <i>Official</i>		84	<i>Terrinthine du Melise</i>		717
<i>Tannojon</i>		88	<i>Terpentanol</i>		1198
<i>Tanno guaraforn</i>		557	<i>Terpin Hydrate</i>	3 to 10 gr	1196
<i>Tannone</i>	5 to 15 gr	88	<i>Terpina</i>		1196
<i>Tannosal</i>		455	<i>Terpineol</i>	2 min	1197
<i>Tanocol</i>	5 to 15 gr	88	<i>Terpinol</i>		846
<i>Taphosote</i>		455	<i>Terra Japonica</i>		1313
<i>Em</i>		905	<i>Test Solution of Starch</i>		1261
" <i>Capsules</i>		909	<i>Tetanus Antitoxin</i>		882
" <i>Oil of</i>		909	<i>Tetradodo phenolphthalein</i>		665
" <i>Light</i>		909	<i>Tetradodo pyrrol</i>		774
" <i>Ointment</i>		909	<i>Tetramethylthionine Hydrochloride</i>		491
" <i>Pills</i>		909	<i>Tetrantrin</i>	10 to 20 gr	1175
" <i>Water</i>		909	<i>Tetronal</i>	3 to 8 gr	1202
<i>Taraxaco</i>		1193	<i>Thallina Sulphas</i>		272
<i>Emulsi Radix</i>		1193	<i>Theobroma Solution Tablet</i>		1207
<i>Emulsi Lmetic</i>		173	" <i>Excipient</i>		1203
<i>Tartarated Antimony</i>		173	<i>Theobromatis Oleum</i>	5 to 10 gr	1204
" <i>Iron</i>		637	<i>Theobromina</i>		1204
" <i>Soda</i>		1088	<i>Theobromine</i>		1205
" <i>Powder, Effervescent</i>		1089	" <i>Sacrylate</i>		1205
<i>Tartarato de Potassi</i>		964	" <i>Sodium Citrate</i>		1206
<i>Lactalic Acid</i>		89	<i>Theocin</i>	3 to 6 gr	1206
<i>Lactaro Emulsi</i>		173	" <i>Sodium Acetate</i>	4 gr	1206
<i>Lactarus Boazatus</i>		968	<i>Theophylline</i>		1206
" <i>Depuratus</i>		966	<i>Theapeutic Agents of Microbial Origin</i>		1260
" <i>Natronatus</i>		1088	" <i>Sera</i>		1260
" <i>Subiatus</i>		174	<i>Thermol</i>		1123
<i>Tartarus Ferrus Potassius</i>		537	<i>Thermonetric Memoranda</i>		cv
" <i>Kalico natrus</i>		1088	<i>Scales Compared</i>		cv
" " <i>Subius</i>		174	<i>Thiocol</i>	10 to 20 gr	687
" <i>Kalurus</i>		964	<i>Thioform</i>		258
" " <i>Acidus</i>		906	<i>Thiol</i>		658
" <i>Natrico Kalurus</i>		1088	<i>Thio oxydiphenylamine</i>		37
" <i>Stribo Kalurus</i>		174	<i>Thiosinamin</i>		1085
<i>Tartrate Loco Potassique</i>		968	<i>Thompson's Fluid</i>		261
" <i>Droit de Potassium et de Sodium</i>		1088	<i>Thonium</i>		1223
" <i>de Morphine</i>		785	" <i>Lactate</i>		1223
" <i>de Potasse Acide</i>		966	" <i>Nitrate</i>		1223
" " <i>Neutrie</i>		964	" <i>Salicylate</i>		1223
<i>Tartrato Antimonico Potassico</i>		173	<i>Thonoughwort</i>		501
" <i>Borico Potassico</i>		908	<i>Thus Americanum</i>		1207
" <i>Ferrus Potassico</i>		537	<i>Thymus Oleum</i>	1 to 2 gr	1210
" <i>de Morfina</i>		788	<i>Thymol</i>		1207
" <i>de Potasio Acido</i>		966	<i>Thymol Antiseptic Dressings</i>		1210
" " <i>Neutro</i>		964	" <i>Camphor</i>		813
" " <i>y Ferro</i>		537	" <i>Carbonate</i>		1211
" " <i>y Sodio</i>		1088	<i>Thymolis Iodidum</i>		1208
" <i>Potassa e de Ferro</i>		537	<i>Thyrodoctin</i>		1214
" <i>Potasse de Soda</i>		1088	<i>Thyroglandin</i>	1 to 5 gr	1214
" <i>Potassio y antimonio</i>		174	<i>Thyroidea</i>		1213
" <i>Sodico Potassico</i>		1088	<i>Thyroidium Siccum</i>	3 to 10 gr	1212
<i>Taraxoholic Acid</i>		1136	<i>Thyroidin</i>		1214
<i>Tincture d Aconit</i>		94	<i>Tinctura Absinthii</i>	1 to 4 fl drms	1
" <i>Balsamique</i>		243	" " <i>Composita</i>		1

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Tinctura Myrrhe et Boracis		263	Tincture of Cudbear		873
" Nuxi Vomica	5 to 15 min	921	" " Myrrh and Borax		885
" Oleri Corticus	30 to 60 min	1059	" " Phosphorus		885
" Opi	{ 5 to 15 min (repeated) 20 to 30 min (single)	833	" " Rhatany		715
" Opi Ammonia	1 to 1 fl drim	841	" " Steel		524
" " Ben oica		310	Tinctures Processes for		1215
" " Camphorata		910	" innelly Senna		1073
" " Citrata		842	Tinospora		1216
" " Decolorati		839	Tintura d Aconito		94
" " Decolorati	8 min	843	" " Alcoholica de Aconito		94
" di Opio Alcanforado		810	" " " Corteca de		391
" " Composita		810	" " Nua nya Compueto		712
" " Persiana		873	" " " Alcoholica de Kola		
" " Phosphori Composita	3 to 15 min	985	" " " Opi Jabo		838
" " Physostigmatis	5 to 15 min	588	" " " nosa		390
" " Phytolacere	3 to 10 min	892	" " " Quina		240
" " Pterorhiza	1 to 1 fl drim	892	" " " Cabalongas		659
" " Podophylli	5 to 15 min	926	" " " di China		390
" " " Ammonia	10 to 30 min	926	" " " China		390
" " " Indica	5 to 15 min	926	" " Chloroformica de Canta		320
" " Plum Virginiane	1 to 1 fl drim	970	" " " rudas		391
" " Pulsatilla	5 to 30 min	972	" " " conoborante de Whitt		1054
" " Pyrethri		973	" " " de Jabor Alcanforado		508
" " " Florum		974	" " " de Mulato de Feio		390
" " Quassia	1 to 1 fl drim	978	" " " de Quina		391
" " Quebracho	1 to 1 fl drim	979	" " " Compuesta		670
" " Quillam	1 to 1 fl drim	980	Tisanes		659
" " de Quina Comp		391	Tisane de Carraquehen		859
" " Quinine	1 to 1 fl drim	994	" " " Gentiane		562
" " " Ammonia	1 to 1 fl dim	1005	" " " Lichen d Islande		859
" " Rhei		1015	" " " d Oranger		225
" " " Amara		1016	" " " d Orge		601
" " " Aquosa		1017	" " " de Polygala		1071
" " " Aromatica		1016	" " " Quinquina		389
" " " Composita	{ 1 to 1 fl drim (repeated) 2 to 4 fl drim (single)	1015	" " " Katanha		714
" " " Dareil		1015	" " " Rhubarbe		1014
" " " Vinosa		1015	" " " Sarsaparelle		1057
" " " Rhus	1 to 5 min	1018	" " " Uva Ursi		1227
" " " Rusa		242	Tobacco Juice		1189
" " " Sabina	20 to 60 min	1081	" " Leaf		1189
" " " Scilla	5 to 15 min	1085	Toddalia		1216
" " " Senega	1 to 1 fl drim	1071	Tolu, Balsam of		217
" " " Senna Composita	{ 1 to 1 fl drim (repeated) 2 to 4 fl drim (single)	1078	Tolu balsam wrop		220
" " " Serpentina	1 to 1 fl drim	1078	Tolypurn	5 to 20 grains	881
" " " Stamonii	5 to 15 min	1165	" " " Salicylate		881
" " " Strophanthi	5 to 15 min	1164	Tolysal		881
" " " Strychni		821	Tropique a l Acetate de Cuivre		465
" " " Succi	25 min	1172	" (Vel)		188
" " " Sumbul	1 to 1 fl dim	1182	Toughened Caustic		1216
" " " Thebaica		838	" "ragac intha		240
" " " Benzoua		310	Traumatic Balsam		589
" " " Thebaicum Crocatum		843	Traumaticine		240
" " " Tinospora	30 to 60 min	1216	Traumatol		44
" " " Lolitana	1 to 1 fl dim	220	Trementina de Alerce		717
" " " Urginea		1065	" " di Venezia		717
" " " Valeriana	1 to 2 fl drim	1229	Tribromo resorcin		1010
" " " Ammonia	1 to 1 fl drim	1228	Tribomphenol		86
" " " Atheria		1229	Tribu hion acetic Acid		14
" " " Indica Ammoni	1 to 1 fl dim	1229	Tribu chlor tertiary butyl alcohol		369
" " " ata		1229	Trichloro methane		871
" " " Vanda		1231	Trichlorphenol		871
" " " Leatri Viridis	5 to 20 min	1231	Triferrin		773
" " " Warburgi		989	Trifoglio Fibroso		778
" " " Whittii		391	Trifolio Fibroso		1218
" " " Zingiberis	1 to 1 fl drim	1257	Trifolium		45
" " " Fortior	5 to 20 min	1257	" " Fibrum		773
" " " "		1215	Trihydroxybenzoic Acid		400
Tincture of Barl. Hualam s		391	Triiodometacis esol		660
			Triiodomethane		42
			Trisresol		1218
			Trimethylamina		
			Trimethylaminæ Hydrochlori	1 to 5 gr	1219
			Trimethylbenzoylpyperidinum		415
			hydrochloricum		

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Unguentum Cicosoti		454	Unguentum Piccolozini		892
" <i>Oreta</i>		457	" <i>Plumbi</i>		921
" <i>Cupri Oleatis</i>		468	" " <i>Acetatis</i>		913
" <i>Diachylon</i>		918	" " <i>Acetici</i>		921
" " <i>Carbolisatum</i>		918	" " <i>Carbonatis</i>		914
" " <i>Hebrae</i>		918	" " <i>Iodidi</i>		910
" <i>Elemi</i>		482	" " <i>Oleatis</i>		918
" <i>Encarnativo</i>		918	" " <i>Oxydatis</i>		918
" <i>Eberna</i>		891	" " <i>Tannici</i>		922
" <i>Eucalypti</i>		498	" <i>Plumbi Baccæ</i>		921
" <i>Gambæ Compositum</i>		550	" <i>Potassæ Sulphuratis</i>		933
" <i>Gallæ</i>		551	" <i>Potassæ Iodidi</i>		950
" " <i>c Opi</i>		551	" <i>Pyrogallol Co</i>		73
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